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DOCTORAL DISSERTATION

Geochemical Characterization of Endmember Mantle Components

by

Rhea K. Workman

June 2005

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Geochemical Characterization of Endmember Mantle Components

By

RHEAK. WORKMAN

B.S., University of Missouri – Rolla, 1998

Submitted in partial fulfillment of the requirements for the degree of

Doctor of Philosophy

at the
MASSACHUSETTS INSTITUTE OF TECHNOLOGY
and the
WOODS HOLE OCEANOGRAPHIC INSTITUTION

June 2005

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Geochemical Characterization of Endmember Mantle Components

by

RHEAK. WORKMAN

Submitted in partial fulfillment for the requirements for the degree of Doctor of Philosophy at the Massachusetts Institute of Technology and the Woods Hole oceanographic Institution

June 2005

Abstract

This thesis uses trace elements and radiogenic isotope tracers to define elemental abundances in reservoirs of the Earth's mantle, including EM2 (the Enriched Mantle 2), as seen in the Samoan hotspot track, and DMM (the depleted upper mantle), which is sampled at mid-ocean ridges. Together these components comprise up to ~50% of the total mantle mass. Much of the mantle's chemical heterogeneities are suspected to originate by either the removal of mass from the mantle (in the case of DMM) or the addition of mass to the mantle through subduction zones (in the case of EM2). We show that DMM represents mantle that 1) has been previously depleted by 2-3% melt removal, 2) mass-balances well with the continental crust, 3) has only 15% of the radiogenic heat production in primitive upper mantle and 4) can generate present-day ocean crust by 6% aggregated fractional melting. EM2 is classically interpreted as mantle material enriched in trace elements through the ancient, subduction-zone recycling of terrigenous sediments; here we show this model is unlikely and provide two other working hypotheses. The first is recycling of meltimpregnated oceanic lithosphere; the second is recycling of a mantle wedge impregnated with melt from a subducting oceanic plate.

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To Boa, Patty, Marcie, Adam and Michael

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Chapter 1:

Introduction

The field of mantle geochemistry is fundamentally concerned with the processes and history of Earth's differentiation. How has Earth developed its inner and outer cores, mantle, continental crust, oceanic crust, ocean and atmosphere? Can we track the growth and evolution of these reservoirs through the 4.56 billion years of Earth history – and can we project into the future? Though we are concerned with all these reservoirs, our field is self-named 'mantle' geochemistry since the Earth's mantle is the primary driver of differentiation. It mediates convective transfer of materials from deep to shallow Earth, resulting in the formation of crust, oceans and the atmosphere. It also returns bits of shallow, differentiated materials back to the deep Earth in places where the mantle is downwelling at subduction zones. The mantle essentially turns the Earth inside-out, trying to purge it of unwanted, incompatible elements (such as hydrogen, carbon, potassium and uranium) and retain in it the wanted, refractory elements (such as magnesium, iron and chromium). It is fascinating to think our human bodies are made of carbon once contained within the very deep, very hot inner Earth.

The Earth was born of materials condensed from our solar system about 4.56 billion years ago (Patterson, 1956; Birck and Allegre, 1978); it almost certainly had some small initial variations in chemical composition, but is thought to be, on average, similar in elemental and isotopic abundances to the class of meteorites called CI carbonaceous chondrites (Anders and Ebihara, 1982; Anders and Grevesse, 1989). From its mess of agglomerated materials, Earth formed (very early in its history) a dense, Fe-Ni metallic core, leaving the remaining lithophile materials to form the Bulk Silicate Earth (BSE); BSE is about 84% the volume of the Earth, but only 68% the mass of the Earth because its density is lower than that of the core. Knowing the approximate mass and composition of the core from seismic surveys through the Earth allows for a fairly good estimate of the chemical composition of the Bulk Silicate Earth (by subtracting the core from the CI composition; Hart and Zindler, 1986; McDonough and Sun, 1995); this simple estimate is inaccurate for

those elements, such as hydrogen, helium, sulfur and lead, that are strongly volatile and lost to space before the formation of Earth's atmosphere.

The BSE has further differentiated, or fractionated, into distinct chemical reservoirs some of which are obvious at the surface of the Earth (like the oceans and continental crust) and some of which are held within the Earth's mantle. The focus of this thesis is on the compositional variations within the mantle, as determined from mantle melts erupted at both mid-ocean ridges and hotspots (i.e. mantle plumes). Zindler and Hart (1986) noted that there are 4 unique end-member mantle components based on 87Sr/86Sr, 143Nd/144Nd, ²⁰⁶Pb/²⁰⁴Pb, ²⁰⁷Pb/²⁰⁴Pb, and ²⁰⁸Pb/²⁰⁴Pb radiogenic isotope compositions of a global database of oceanic lavas. They described how most mid-ocean ridge basalts (MORBs) are isotopically similar and represent an upper mantle previously depleted of melt (i.e. oceanic and continental crust); this reservoir is termed the Depleted MORB Mantle (DMM). On the other hand, ocean island basalts (OIBs), erupted by hotspot volcanism, are isotopically heterogeneous in terms of most radiogenic isotope systems. Each hotspot chain is unique in radiogenic isotope space and represents a unique mixture of the Zindler and Hart mantle components. From parent isotope half-lives and parent-daughter ratios, it is inferred that end-member mantle sources for OIBs and MORBs must have been chemically isolated for billions of years in order to develop the observed differences in the abundance of daughter isotopes. Because isotopes of heavy elements are so little fractionated in the melting process, isotopic compositions of oceanic basalt are not only "clocks" for ancient reservoir development, but also "fingerprints" of a melt's solid source. The goal of mantle geochemistry in general and this thesis specifically is to determine the origins, ages and chemical compositions of these mantle components.

Although there have been many ideas regarding the origins of the classic mantle endmembers seen in OIB chains, one model has been relied upon most commonly and received
the most attention from a modeling point of view. We are in effect "outside looking in", so
major differentiation processes occurring at the solid Earth's uppermost layers, namely the
formation of continental and oceanic crust, are the most obvious explanations for the
creation of volumetrically significant heterogeneities in composition. Return of these
differentiated materials to the mantle through return flow at subduction zones creates *mantle*heterogeneity. Geochemical models attempting to accurately quantify the compositions of
deeply-subducted materials need precise knowledge regarding: 1) hydrothermal alteration of
the oceanic crust, 2) partition coefficients for both the dehydration and melting of crust and
sediments, 3) the thermal structure of mantle wedges, 4) the compositions of subducting

sediments, and 5) the lifespan of a subducted slab in the deep mantle. Although much progress has been made in each of these topics, many parameters are not well defined through space and time, making it difficult to formulate a completely accurate model for any one of the OIB end-members.

In this thesis, I focus on describing the chemical character of 2 out of the 4 mantle reservoirs that were identified and 'fingerprinted' by Zindler and Hart (1986). The first is DMM; it is ubiquitously sampled along the global system of mid-ocean ridges and represents the largest accessible reservoir on Earth (at least 30% the BSE). It is called 'depleted' because we know from its isotopic fingerprint that melts were removed from this mantle a very long time ago (2 to 3 Ga), in a process we speculate to be much like the oceanic and continental crust formation that occurs today. We use the following constraints in deriving an average trace element composition of DMM: 1) trace element content of clinopyroxenes from abyssal peridotites, 2) isotopic evolution from primitive upper mantle (PUM) and 3) canonical trace element ratios in MORBs. The average trace element content of DMM, as deduced here, generally shows a very smooth pattern with increasing trace element compatibility, which to first order mass-balances with the continental crust. The degree of depletion indicated in DMM represents 2-3% melt removal from the primitive upper mantle; this means that DMM has only 15% the radiogenic heat production of PUM (from K, U and Th). Present-day ocean crust (i.e. average MORB) can be modeled with 6% aggregated fractional melting of the deduced DMM.

The second mantle reservoir studied in this thesis is nearly opposite in character from the depleted upper mantle – it is called the Enriched Mantle 2 (EM2) and is sampled from the inner Earth by the Samoan hotspot. EM2 has the highest ⁸⁷Sr/⁸⁶Sr of all oceanic lavas, whereas the depleted upper mantle has the lowest. Although EM2 may not be a volumetrically significant reservoir, its rare and unique composition is very significant because it allows us to eventually understand the details of BSE differentiation (for example, the growth rate of continental crust).

Chapters 2, 4 and 5 of this thesis address the geochemistry of the Samoan islands and seamounts in great detail. In Chapter 2, we use trace element abundances and radiogenic isotope tracers to establish the geochemical variability, character, and habits of the Samoan lavas, including the ones that are more extreme in their EM2 signature than any other lavas sampled on Earth. In Chapter 2, we establish that a model involving the recycling of only marine sediments will not successfully explain the origin of EM2. Instead, an alternative model is offered in which an oceanic lithosphere is impregnated with an upper mantle melt,

recycled through a subduction zone and stored for 2.5 Ga in the mantle – to be sampled today by the Samoan mantle plume.

In Chapter 4, we use volatile element abundances (water, carbon dioxide, fluorine, sulfur and chlorine) from Samoan glasses to comment on the general properties of these elements in igneous systems and the specific properties of water in the EM2 mantle reservoir. We find that absolute water contents are high for Samoan lavas (0.63 – 1.50 wt%), but relative enrichment of water compared to trace element enrichment is low. H₂O/Ce (58 – 157) and H₂O/La (120 – 350) correlate inversely with ⁸⁷Sr/⁸⁶Sr compositions (0.7045 – 0.7089). This leads us to believe that, because of very fast diffusion of hydrogen in olivine, recycled lithospheric material with high initial water content will lose water to the drier ambient mantle during storage within the inner Earth. This concept implies that water may be one of the few (if only) elements in the mantle that is close to chemical equilibrium over great distances.

Chapter 5 presents oxygen isotope data for Samoan olivine phenocrysts as a means to identify the presence of recycled material that was once at or near Earth's; oxygen isotope variations occur not because of production of a radiogenic oxygen isotope, but because oxygen's isotopes, ¹⁶O and ¹⁸O, have slightly different chemical behavior during low temperature reactions. Here we test two models for the generation of EM2. The first is (once-again) the standard sediment-recycling model. The second expands upon the metasomatic model of Chapter 2 by identifying a location for the metasomatism (the mantle wedge), and more accurately defining what the metasomatising agent is (an eclogite melt). Each model shows some major misfits to characteristics of end-member Samoan lavas, although the metasomatic model requires less 'special pleading'.

Chapter 6 investigates the correlation between geochemical and geophysical properties of Samoa. Excess crustal flux along the Samoan volcanic lineament decreases nearly monotonically approaching Vailulu'u Seamount, the easternmost and youngest volcano. This trend shows excellent correspondence to increasing ²⁰⁶Pb/²⁰⁴Pb compositions of the lavas with decreasing age along the hotspot track. We speculate this correlation could be due to either a lithosphere thickening toward the east or decreasing potential temperature of the mantle plume, and that the geochemical signatures observed in the lavas are partly a function of how mantle materials with differing solidus temperatures are sampled from the Earth.

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Chapter 2:

Recycled metasomatized lithosphere as the origin of the Enriched Mantle II (EM2) Endmember: Evidence from the Samoan Volcanic Chain*

Abstract

An in-depth Sr-Nd-Pb-He-Os isotope and trace element study of the EMII-defining Samoan hotspot lavas leads to a new working hypothesis for the origin of this high ⁸⁷Sr/⁸⁶Sr mantle endmember. Systematics of the Samoan fingerprint include 1) increasing ²⁰⁶Pb/²⁰⁴Pb with time – from 18.6 at the older, western volcanoes to 19.4 at the present day hotspot center, Vailulu'u Seamount, 2) en-echelon arrays in ²⁰⁶Pb/²⁰⁴Pb – ²⁰⁸Pb/²⁰⁴Pb space which correspond to the two topographic lineaments of the 375 km long volcanic chain this is much like the Kea and Loa Trends in Hawai'i, 3) the highest 87Sr/86Sr (0.7089) of all oceanic basalts, 4) an asymptotic decrease in ³He/⁴He from 24R_A (Farley et al., 1992) to the MORB value of 8R_A with increasing ⁸⁷Sr/⁸⁶Sr, and 5) mixing among four components which are best described as the "enriched mantle", the depleted FOZO mantle, the (even more depleted) MORB Mantle, and a mild HIMU (high ²³⁸U/²⁰⁴Pb) mantle component. A theoretical, "pure" EMII lava composition has been calculated and indicates an extremely smooth trace element pattern of this endmember mantle reservoir. The standard recycling model (of ocean crust/sediment) fails as an explanation for producing Samoan EM2, due to these smooth spidergrams for EM2 lavas, low ¹⁸⁷Os/¹⁸⁸Os ratios and high ³He/⁴He (>8R_A). Instead, the origin of EM2 has been modeled with the ancient formation of metasomatised oceanic lithosphere, followed by storage in the deep mantle and return to the surface in the Samoan plume.

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1. Introduction

Although intra-plate ocean island volcanism accounts for only a few percent of the total volcanism on Earth, these volcanic piles may be the surface manifestations of the deepest known samplings of the interior of the planet. The relative stationarity of mantle plumes with respect to upper mantle plate flow (Molnar and Stock, 1987; Steinberger and O'Connell, 1998; Wang and Wang, 2001; Koppers et al., 2001), and a growing catalogue of seismic evidence and tomographic images showing velocity anomalies beneath hot spots extending well into the mid-mantle and sometimes to the core-mantle boundary (Russell et al., 1998; Shen et al., 1998; Zhao, 2001; Montelli et al., 2003), all support the idea that mantle plumes sample the inner Earth at a much deeper level than do mid-ocean ridge spreading centers. Ocean island chains may thus provide some of the best clues to the chemical character of the lower mantle and the nature of convective interactions between the deep and shallow mantle.

Unlike mid-ocean ridge basalts (MORBs), which derive from a fairly uniform meltdepleted upper mantle, ocean island basalts (OIBs) are isotopically heterogeneous in terms of most radiogenic isotope systems (e.g. Zindler and Hart, 1986; Hart, 1988; Hofmann, 1997). Isotopic arrays from ocean island chains often extend from a "common" mantle, termed FOZO (i.e. Focus Zone; Hart et al., 1992), and tend toward one of three "endmember" mantle components: HIMU, the high time-integrated U/Pb mantle, EM1 or EM2, the Enriched Mantles 1 and 2 (Zindler and Hart, 1986). From parent isotope half-lives and parent-daughter ratios, it is inferred that mantle sources for OIBs and MORBs must have been chemically isolated for billions of years in order to develop the observed differences in the abundance of daughter isotopes. Because isotopes of heavy elements are so little fractionated in the melting process, isotopic compositions of oceanic basalt are not only "clocks" for ancient reservoir development, but also "fingerprints" of a melt's solid source. We are left, through geochemical interrogation and theoretical ingenuity, to reverse the processes by which mantle melts were generated and brought to Earth's surface. Ultimately, with some indication for source compositions, the origins and ages of chemically distinct, isolated mantle reservoirs can be deduced.

Although there have been many ideas regarding the origins of the classic mantle end members, one model has been relied upon most commonly and received the most attention from a modeling point of view. We are in effect "outside looking in", so major differentiation processes occurring at the solid Earth's uppermost layers, namely the formation of continental and oceanic crust, are the most obvious explanations for the creation of volumetrically significant heterogeneities in composition. Many workers have applied this perspective and contributed to what is here referred to as the Standard Model for the origin of mantle components (Armstrong, 1968; Chase, 1981; Hofmann and White, 1982; Cohen and O'Nions, 1982; White, 1985; Zindler and Hart, 1986; Weaver, 1991; Hart et al., 1992). In summary, oceanic crust is subducted at convergent margins, dehydrated (increasing U/Pb, Th/Pb, and Sr/Rb ratios) and put into long term storage in the deep mantle to evolve to HIMU. EM1 and EM2 are generated when trace-element-enriched pelagic (i.e. deep-sea) and terrigenous (i.e. continental) sediment, respectively, accompany the subducted and stored oceanic crust (Fig. 1). Geochemical models attempting to accurately quantify the compositions of these deeply-subducted materials (Hart and Staudigel, 1989; Weaver, 1991; Stracke et al., 2003) are greatly hindered by a lack of knowledge regarding: 1) hydrothermal alteration of the oceanic crust, 2) partition coefficients for both the dehydration of crust and sediments and the melting of sediments, 3) the thermal structure of mantle wedges, 4) the variable compositions of sediments in space and time, and 5) the lifespan of a subducted slab in the deep mantle. Although much progress has been made in each of these topics, the constraints are not strong enough to provide the needed resolution in parent/daughter ratios. Ironically, it may be exactly the lack of constraints that ultimately makes the Standard Model nonviable. By all indications from today's geodynamical systems, sediments and the subduction zone processing of crust and sediments all display such variability that a specific composition (which evolves to HIMU, EM1 or EM2) almost certainly would not be produced twice, and there would be no discrete or recognizable "endmember" reservoirs. On the other hand, and often the strongest criticism of the Standard Model (e.g. Hawkesworth et al., 1984; Barling and Goldstein, 1990; Morgan, 2000), is that there may be no such things as mantle endmembers. Each ocean island array could consist of its own unique isotopic composition, which represents a unique subducted slab from a unique recycling time.

In the present study, we specifically deal with the origin of the Enriched Mantle II (EM2) endmember. Lavas from the Samoan Islands have long been recognized as holding the most extreme signal of EM2 (Zindler and Hart, 1986; Wright and White, 1986; Farley et al., 1992, Hauri and Hart, 1993). Here we use a new comprehensive geochemical study to assess possible origins of the EM2 reservoir. This paper outlines why the recycling of sediment/slab cannot be the origin of EM2, and offers an alternative model which will

generally result in consistent trace element compositions, and hence isotopic signatures, through time. We assume that mantle endmembers do, in fact, exist, and that one process, acting to varying degrees at a variety of times, will produce a fairly homogeneous endmember reservoir, which is available for mixing with other mantle components during upwelling of mantle plumes.

The working model introduced here for the origin of EM2 involves metasomatism (i.e. fluid/melt infiltration) of oceanic lithosphere, followed by subduction zone recycling and long-term storage of this lithosphere. As a process for creating trace-element-enriched mantle, metasomatism is not a new idea and has been invoked both for continental lithosphere (Frey and Green, 1974; Brooks et al., 1976; Menzies and Murthy, 1980; Menzies, 1983) and oceanic lithosphere (Zindler et al., 1979; Kay, 1979; Hawkesworth et al., 1979, 1984; Richardson et al., 1982; Roden et al., 1984; Hart et al., 1986; Halliday et al., 1992; Class and Goldstein, 1997; Niu et al., 1996; 1999; Niu and O'Hara, 2003). The process we envision is much like the SYS model of Zindler et al. (1979), and the autometasomatic model of Roden et al. (1984). We envision it operating on newly formed lithosphere close to spreading centers, as illustrated by Niu et al. (1999) and Niu and O'Hara (2003).

We show that a lithosphere impregnated 2.5 Ga with a small-degree upper mantle melt can evolve to the present day isotopic composition of EM2. This model provides an EM reservoir with much greater volume than that of oceanic crust and sediment. A more voluminous "package" will have greater resistance to mixing within the convecting mantle and therefore have greater possibility of staying an isolated body for the required 2.5 Ga evolution time. Another benefit of this model is that the lithosphere will be isolated and protected from subduction zone processing (such as elemental fractionations that occur within the subducted oceanic crust and sediments during metamorphism and devolatilization).

2. Geologic setting

The Samoan islands and seamounts are centered on 14°S latitude and stretch from 169-173°W longitude (Fig. 2). They sit ~100 km north of the northern termination of the Tonga Trench, on ~110 Ma oceanic crust of the Pacific Plate which is moving 25.8° WNW at 7 cm/yr (Sella et al., 2002). The Samoan volcanoes separate into two topographic ridges, both sub-parallel to the direction of plate motion: the Savai'i – Upolu – Tutuila – Malumalu

group define the southwestern (and generally older) lineament, and the Muli – Ofu/Olosega – Ta'u – Vailulu'u group define the northeastern (and younger) lineament. We will designate these the "Malu" and "Vai" Trends, respectively. The recently mapped leading-edge seamount, Vailulu'u, rises from 5000 meter seafloor to a summit depth of 590m (Hart et al., 2000). Recent volcanic activity at Vailulu'u has been documented with the following observations: elevated water temperatures and particulate contents within the summit crater, a halo of intense particulate matter surrounding the summit in the depth range of 600-800 meters, high Mn concentrations and ³He/⁴He ratios (up to 9 R_A) in the crater water, swarms of seismicity, and dredged rock samples with U-series ages of 5-50 years (Hart et al., 2000). The age-progression heading west from this present-day hot spot location approximately follows the plate velocity of 7 cm/yr and includes the seamounts Lalla Rookh, Combe, and Alexa, which is 1750 km west of Vailulu'u (Duncan, 1985; Natland and Turner, 1985; McDougall, 1985; Hart et al., 2000; Hart et al., unpubl. data). Malulu seamount and Rose Atoll to the east of Vailulu'u do not have Samoan isotopic signatures (Hart et al., unpubl. data), and are most likely associated with the Cook-Austral lineament.

As if burning the candle at both ends, post-erosional volcanism has been extensive on the westernmost island of Savai'i (with the most recent eruptive episode taking place from 1905-1911) as well as being documented on the islands of Upolu and Tutuila (but here, all pre-historic, and much less extensive) (Kear and Wood, 1959; Keating, 1992). Although the pervasive post-erosional veneer on Savaii has disrupted the age-progression model (Savai'i should be ~5Ma based on the plate velocity model) and has lead to debates about the origin of the Samoan volcanoes (e.g. Natland, 1980), we believe there is little doubt about the chain originating from hot spot volcanism. The atypical volume of post-erosional volcanism on Savai'i is possibly due to the complicated tectonic setting of the volcanic chain. Since Savai'i is closest to the Tonga Trench, it is reasonable that bending stresses are facilitating additional melt extraction from the upper mantle (e.g. Hawkins and Natland, 1975; Natland, submitted 2003).

Tectonic reconstruction of the region (Brocher, 1985; Pelletier et al., 1998; Zellmer and Taylor, 2001) show that the transform-fault bounding the northern Tonga Trench evolved ~6-8 million years ago from the fossil Vitiaz Trench in response to opening of the Lau back-arc basin. Studies of the chemical characteristics of the northern Lau back-arc basin seamounts and seismic profiling beneath the basin collectively suggest leakage of Samoan plume material into the northern Lau Basin through a tear, or window, in the paleo-slab of the Pacific Plate subducted at the Vitiaz Trench. Geochemical evidence includes high

³He/⁴He lavas of some Lau Basin seamounts (Poreda and Craig, 1992; Turner and Hawksworth, 1998), with trace element and isotopic compositions which are more characteristic of OIBs than MORBs or IABs (Ewart et al, 1998; Danyushevsky et al, 1995; Wendt et al., 1997). Seismic studies by Millen and Hamburger (1998) and Chen and Brudzinski (2001) illustrate a remnant slab of the Vitiaz subduction that has detached from the warped Pacific Plate, thereby providing an unobstructed path for melt/mantle migration from the Samoan plume into the Lau Basin. By speculation, this suggests that the Samoan plume beneath the Pacific Plate is much more widespread than the discrete lineament of volcanoes would indicate. Also, the exact location of the Samoan volcanoes may not necessarily be where the plume upwelling is "strongest", but instead where the plume fortuitously intersects a structural weakness imparted to the lithosphere by tectonic stresses of the local area. The en-echelon nature of the volcanic edifices may provide witness to this structural control (see Natland, submitted 2003, for a full discussion of this idea).

3. Samples and Analytical Details

3.1. Sample locations and descriptions

Rock samples utilized in this study have been collected from both land and sea. The seamounts Vailulu'u, Muli, and Malumalu, along with submarine portions of Ta'u, were dredged during the 1999 AVON2/3 cruise of the R/V Melville. Land-based sampling of Savai'i and Upolu, conducted in 2001, was aimed at expanding the coverage of "old shield" (namely, the Fagaloa Volcanic Series; Kear and Wood, 1959), and thereby establishing a greater temporal coverage of the Samoan plume. On Upolu, we sampled the southwestern exposure of the Fagaloa Volcanics; this is a topographic massif with well-developed river valleys, referred to as A'ana by the local inhabitants. Our Upolu samples primarily come from along or near the Matafa'a coastline and Fagalei Bay. Samples from Savai'i were collected from the north-central shore, where exposures of Fagaloa Volcanics were mapped over a 20 km² relative topographic high (Kear and Wood, 1959). This area is bound to the east by the village of Vaipouli, contains the Muliolo and Eatelele Streams, and is bound to the west by an escarpment that leads down to the village Paia.

Subaerial sampling of Ta'u, the youngest island of the chain, was conducted in 1999 and was principally concentrated along the coastline. The sampling was temporally diverse, in that all five of the volcanic series mapped by Stice and McCoy (1968) are represented.

Unlike the older and larger islands of Savai'i and Upolu, Ta'u Island manifests from only one main shield volcano; this simplified structure is reflected in the isotopic homogeneity observed for Ta'u, as will be discussed in following sections.

Phenocryst abundances in Samoan lavas range from 0% to 50% and include the following minerals in decreasing modal abundance: olivine, clinopyroxene, plagioclase, orthopyroxene, and Ti-augite. Phenocrysts are most common in samples from Vailulu'u and least common in samples from Savai'i and Upolu. In thin section, some samples show two populations of olivine in which a coarse-grained population (2-10 mm) shows resorption boundaries and a smaller-grained population (1-2 mm) shows almost no embayed crystal boundaries. However, for most samples, olivine major element compositions (Jackson et al., unpubl.) show that phenocrystic olivines are in Mg-Fe equilibrium with the coexisting liquids. Some samples (especially T14) have glomerocrysts of olivine (±spinel). Plagioclase, clinopyroxene, and oxides are the most common matrix minerals. Hand-samples can generally be classified as aphanitic basalt, olivine basalt, picrite or (rarely) ankaramite. Alteration, in the form of iron-oxide, is most prevalent in the Savai'i and Upolu samples. Sample 63-11 from Vailulu'u crater shows hydrothermally-precipitated quartz rinds along some cracks and grain boundaries.

3.2. Analytical techniques

Techniques reported here are for samples described above. Additional subaerial samples from Savai'i, Upolu, Tutuila and Ta'u have been collected by K.A. Farley and J.H. Natland over the last two decades and analyzed by K.A. Farley for Sr-Nd-Pb-He isotopic compositions. Additional subaerial samples from Savai'i and Upolu have been collected and analysed for major and trace elements and Sr-Nd-Pb isotopes by M. Regelous. We include these data in the present manuscript, as they are previously unpublished; any differences in analytical techniques are reported in the corresponding data tables.

Sr, Nd, and Pb isotopic analyses were carried out with conventional ion exchange procedures (references in Taras and Hart, 1987), using whole rock powders, prepared in an agate shatterbox, and leached for 1 hour in warm 6.2 N HCl. The TIMS techniques are described by Hauri and Hart (1993). Sr and Nd isotope data carry 2σ precisions of ±35 ppm and ±40 ppm, and are reported relative to 0.71024 (NBS 987) and 0.511847 (La Jolla), respectively. Some samples run for Sr and Nd by NEPTUNE multi-collector ICP/MS at W.H.O.I. are of comparable precision to TIMS analyses. The precision of TIMS Pb data is taken to be 0.05% per mass unit after fractionation-correcting to the NBS

981 values given by Todt et al. (1996). Pb isotopic compositions of some samples were also determined on the P54 multi-collector ICP/MS in Lyon, with 2σ precisions of all ratios of ~200 ppm. Additionally, the Upolu and Savai'i sample suite was analyzed on the NEPTUNE multi-collector ICP/MS at W.H.O.I.; using a Tl internal standard, the 2σ external reproducibility for these samples was ±100 ppm or better for all ratios (see Hart et al., 2002). Helium isotopic compositions (3He/4He RA, relative to atmospheric standard) of olivine and/or fresh glass separates (~1-3 mm) were determined at W.H.O.I. by in vacuo crushing, using methods described in Kurz et al. (1996). Analytical errors average ±0.2 Ra at 2σ, for helium concentrations ranging from ~10⁻⁸ to 10⁻⁶ cc/gram. Os isotopic compositions on a select group of olivine-rich samples were determined by sparging of OsO4 into W.H.O.I.'s Finnigan Element Magnetic Sector ICP-MS, following a flux fusion sample preparation (see Hassler et al., 2000 for a detailed Os analytical technique). Fusion blank corrections resulted in 0.06-1.22% corrections to the ¹⁸⁷Os/¹⁸⁸Os ratios. Major elements and some trace elements (Ni, Cr, Sc, V, Ga, Cu, Zn) in unleached whole rock powders were measured by XRF, and all other trace elements by ICP/MS at Washington State University (Hooper et al, 1993). Submarine glasses have been analyzed for major elements by electron microprobe at Massachusetts Institute of Technology.

3.3. Sample preservation/quality

Despite sampling of lavas from older shield and submarine settings, the quality of preservation is generally very good. The Th/U ratios of the sample suite fall entirely within 4.5 ± 1.5 (with the exception of sample S15 at Th/U = 6.7) and show a slight (although rough) positive correlation with Th concentrations. The Ba/Rb ratios have an average of 9.3 \pm 1.8 at 1 σ (near the canonical value of ~12 for fresh ocean island basalts; Hofmann & White, 1983) and are inversely correlated with Rb concentrations; significant exceptions to this correlation are samples 79-4, S15, and S25, with Ba/Rb ratios of 17.2, 14.0, and 3.7, respectively. We take these two proxies of alteration as indications that elements as or less mobile than Rb and U are very nearly pristine for most samples. However, elevated Rb/Cs ratios (176 \pm 70 at 1 σ) in the subaerial Upolu and Ta'u samples are most likely explained by chemical weathering and contrast strongly with the roughly canonical values (85-95; Hofmann & White, 1983) represented by the remaining suite (97 \pm 30 at 1 σ).

4. Age Relationships and Age-Progression

Vailulu'u seamount, the most easterly volcano in the Samoan chain, is currently active and belived to be the present day hotspot center (Hart et al., 2000). U-series data constrain two samples from Vailulu'u's summit region to be less than 50 years old; 7 other samples from six dredge locations show excess ²³⁰Th/²³⁸U, evidence of ages less than a few hundred thousand years (Sims and Hart, to be submitted). The oldest K-Ar age from Tau Island is 0.3 Ma (McDougall, 1985). The youngest volcanic series on Tau (Faleasao) is probably younger than 37,000 years, based on ¹⁴C ages of coral inclusions in these volcanics (Hart, unpublished). Additionally, there was an underwater eruption just west of Tau in 1866 (see description in Keating, 1992), evidence that Tau is still in an active shield-building stage. As yet, we have no age constraints on Muli seamount, though the samples dredged from there appear "older" than those dredged from Vailulu'u or Tau. Samples from three dredges on Malumalu show ²³⁰Th/²³⁸U excesses similar to those on Vailulu'u (Sims and Hart, to be submitted), suggesting that Malumalu is not significantly older than Vailulu'u. K-Ar ages for the Pago and Masefau shields on Tutuila range from 1.0-1.9 Ma (McDougall, 1985; Natland and Turner, 1985), somewhat younger than the 2.3-2.7 Ma expected from plate motion considerations.

New high-quality ⁴⁰Ar/³⁹Ar step-release plateau ages are given in Table 1 for the northern shield on Savai'i and the SW shield on Upolu, along with an earlier 40/39 total fusion age for the Vanu River shield on Savai'i. Previous K-Ar ages on the eastern Upolu shield range from 1.54-2.74 Ma (Natland and Turner, 1985); our western shield ages are 0.93 and 2.65 Ma. The older age agrees with the older ages of the eastern shield, though both shields appear younger than the expected plate model age range of 3.9-4.5 Ma. The 0.93 Ma sample (U10) was collected from well within the interior of the eroded SW shield massif, and appears to be reliable evidence for an extended (~2 Ma) period of shield building on Upolu.

There are no published radiometric ages from Savai'i. Based on a plate velocity of 7 cm/year, the age expected for shield initiation on Savai'i is about 5.2 Ma; the two ages reported in Table 1 for the northern (Manase) shield, 0.24 and 0.39 Ma, are far younger than this expected plate age. Kear and Wood (1959) mapped this northern area as shield largely on the basis of abundant surface streamflow. However, we found no obvious evidence of unconformable erosional morphology in this area, and the geochemical evidence discussed below strongly suggests that this map unit is akin to the post-erosional basalts on

Savai'i and Upolu, and unlike the Upolu shield basalts. The young ⁴⁰Ar/³⁹Ar ages are consistent with a re-assignment of this unit to post-erosional status. In the southern interior of Savai'i, Kear and Wood (1959) mapped a small exposure of shield in a gorge on the upper Vanu River. This area is virtually inaccessible, but a trachyte cobble was collected from the lower Vanu River by one of us (KAF) in 1991, and the 40/39 total fusion age of this trachyte is 2.05 Ma (Table 1). While still significantly younger than a plate-model age, this trachyte age is nevertheless very important as it shows that not all of the volcanism on this island can be related to proximity to the Tonga trench, as suggested by Natland (1980); at 2 Ma, the corner of the Tonga trench was almost 400 km west of Savai'i (Bevis et al., 1995). On the other hand, there can be little doubt that Savai'i has been massively resurfaced with post-erosional volcanism as proposed by Natland (1980). The early history of this island will probably only be accessed by dredging on the deeper flanks, where slope failure provides an exposed record.

All in all, the radiometric ages of shield lavas in Samoa are broadly consistent with a simple age-progressive hotspot track, in that ages generally increase from east to west. However, it is clear that shield ages are overall younger than those predicted by plate motion, most likely because the oldest incipient shield lavas are not sampled at the surface of present day volcanoes. While the earliest stages of shield building on Tutuila, Upolu and Savai'i are thus far missing from the sampled record, it would be premature to use this as evidence against a simple hotspot model for Samoa.

5. Magma Generation and Crystal Fractionation

In major element composition, the Samoan basalts and trachybasalts analyzed for this study are clustered just above the alkali-tholeiite line (MacDonald and Katsura, 1964) at 44-49 wt% SiO₂ (Fig. 3; Tables 2 and 5). Samples that fall into the tholeiitic field are, for the most part, from Vailulu'u Seamount, and three of these are highly picritic. Post-erosional lavas (on Savai'i, Upolu and Tutuila) overlap with the shield volcanics, but extend to much greater silica-undersaturation (basanites and nephelinites down to 36 wt% SiO₂; Hawkins and Natland, 1975; Johnson, 1983; Hauri and Hart, 1997). Mg#'s (molar percent Mg/Mg+Fe²⁺) range from 40 in the differentiated Muli samples to 85 in the Vailulu'u picrites. The low MgO/high SiO₂ end of the suite is (vaguely) dominated by samples from Vailulu'u, Malumalu, and Upolu; on the other hand, high MgO/low SiO₂ samples are mainly from Savai'i, Tutuila, and Ta'u. Also plotted on some co-variation plots of Figure 3

are trajectories of near-solidus primary melt compositions at varying pressures of melting in the garnet stability field, using algorithms defined by Herzberg and Zhang (1996) through experiments on KLB-1 peridotite. Recent experiments on another fertile peridotite (KR4003 starting material; Walter, 1998) show primary melt compositions with a general shift to higher MgO at a given pressure (by ~5% in the 4-5 kbar pressure range). Most of the Samoan lavas have undergone some amount of crystal fractionation, as indicated by the fact that they have significantly lower Mg#'s than any estimated primary mantle melts.

The relationship between Mg#'s and CaO shown in Figure 4 provides information regarding both magma generation and crystal fractionation. Plotted along with lava compositions is the trajectory of primary mantle melt compositions (Herzberg and Zhang; 1996). To assess the extent of differentiation and the minerals involved, we have used the pMELTS program (Ghiorso et al., 2002) to model anhydrous fractional crystallization (at a best-fit pressure of 3-4 kbar) of some of the more primitive lavas (Ta'u samples T14 and T48, Vailulu'u sample 63-3, and Malumalu sample 78-1). PMELTS trajectories calculated with 1 wt% H₂O, at a given pressure, are nearly identical to anhydrous runs at 1kbar lower pressure (not shown). All starting compositions have olivine as the second liquidus phase (after spinel), leading to a negative slope for the liquid path on Mg# against CaO (Fig. 4). The kink to positive slopes on the liquid lines of descent marks the crystallization of cpx±olivine±plagioclase. As CaO content of the starting magma decreases, clinopyroxene (cpx) saturation occurs at lower Mg#'s. A suite of submarine Samoan glasses (Table 3) has also been plotted on Figure 4 and shows that true liquids follow the lines of crystal fractionation predicted by pMELTS.

Suites of lavas from each volcano cluster along fairly distinct Mg#-CaO fractionation trends. By projecting the olivine fractionation trends for the different volcanoes back to the primary mantle melt trajectory, we can interpret that the Samoan lavas were generated in the pressure range of 2.5-6 GPa; the order of increasing pressure of melting would be Vailulu'u < Ta'u/Malumalu < Tutuila/Upolu/Savai'i. The extrapolated primary magmas in this model have an extensive range in MgO, from 11 wt% at lowest pressure to 22 wt% at 6 GPa. To get integrated pressures of melting as high as 6 GPa, melting would have to initiate at depths exceeding 180 km and terminate at depths much deeper than the thickness of the lithosphere (~100 km). Given estimates for potential temperatures of plumes (~1550°C; Watson and McKenzie, 1991) and water-undersaturated solidi (Hirth and Kohlstedt, 1996), the depth of *initial* melting is close to 180 km and therefore cannot be the *integrated* depth of melting. The above approach is strictly valid only if the lavas from each

volcano are derived from a constant source composition. We conclude that this CaO index for pressure of melting is rather rickety, given the isotopic variations between volcanoes that will be discussed further into the paper, and that the order of "increasing pressure of melting" is clouded by the extent to which melts were generated from a depleted (low CaO) material.

The Vailulu'u samples are not only high in CaO (also Ca/(Ca+Na) and CaO/Al₂O_{3.}) at a given MgO value, but they are also low in Na₂O, TiO₂, and FeO. This suggests they have the most promise in being interpreted as the shallowest, highest degree partial melts in the whole sample suite (Kinzler and Grove, 1992; Herzberg and Zhang, 1996; Walter, 1998). Melting beneath the other volcanoes may be initiated deeper in the mantle, possibly due to (1) differences in source composition (required by isotopic variations), (2) higher potential temperatures and mantle flow rates, or (3) mantle flow paths which affect melt-solid segregation. The Vailulu'u suite is fit fairly well by a crystal fractionation trend at a pressure of 3 kbar, and indicates cpx fractionation has likely occurred for most samples.

For Ta'u and Malumalu, olivine fractionation clearly dominates the spread in lava compositions. A few samples with Mg#'s greater than ~73 have obviously accumulated olivine (they are phenocryst-rich), but most samples lie along olivine fractionation lines or at the intersection of the olivine control line and cpx saturation (Fig. 4). Three Ta'u samples have compositions close to those of the Muli samples and have surely undergone cpx fractionation; these samples also have the lowest concentrations of the cpx-compatible elements vanadium and scandium in the whole suite (not shown). If parental magmas for all the Ta'u and Muli samples were of nearly the same composition, liquid lines of descent indicate that these low Mg# lavas have undergone about 15% more olivine fractionation than samples T14 and T48, along with 25% cpx fractionation.

6. Isotopes and Trace Elements

6.1 The Global Context

Plotted on the three-dimensional axes of Figure 5 is the mantle tetrahedron of Hart et al. (1992), with data from the ocean island chains which quintessentially define the coordinates for each of the mantle components, EM1, EM2, and HIMU. Data arrays for individual island chains, as well as groups of taxonomically similar island chains, quasi-linearly extend from one of the three OIB endmember components toward FOZO, the common mantle;

very notable is the serious lack of elongation of arrays along tie-lines between the three OIB components. It is clear that EM2 lavas in general, and Samoan lavas in particular, dominate the range in oceanic ⁸⁷Sr/⁸⁶Sr values, but are much less variable in ¹⁴³Nd/¹⁴⁴Nd than EM1. The variation in ²⁰⁶Pb/²⁰⁴Pb found in EM2 basalts is small relative to the composite oceanic suite.

Strontium, neodymium, lead, helium, and osmium isotope ratios for Samoan basalts are given in Tables 4 through 6. Isotope plots (Figs. 6-9) show this new data along with data reported in previous studies (Wright and White, 1986; Farley et al., 1992; Hauri and Hart, 1993). The wide range in ⁸⁷Sr/⁸⁶Sr values, 0.7044 – 0.7089, is correlated with the more narrow range of 0.51293 – 0.51251 for ¹⁴³Nd/¹⁴⁴Nd (Fig. 6). Each island or seamount tends to show a unique field of isotopic compositions that, as will be shown, evolve systematically through space and time. Malumalu Seamount contributes the furthest afield EM2 signature and now defines the most radiogenic ⁸⁷Sr/⁸⁶Sr value (0.7089) of all oceanic lavas. At lower ⁸⁷Sr/⁸⁶Sr (0.7044), near estimates for Bulk Silicate Earth (BSE), the Samoan array is split into two prongs – the "serpent's tongue". Both prongs, one comprised of lavas from Ta'u Island and the other, at higher ¹⁴³Nd/¹⁴⁴Nd, comprised of lavas from Upolu and Tutuila, are significantly elevated (at £_{Nd} of +3 and +5, respectively) over the BSE value of 0.512638 (Hamilton et al., 1983). The other notable EM2 hotspot, the Societies, overlaps the lower prong of the "serpents tongue", and is generally shifted to less-enriched Sr and Nd values. The classic EM1 array (Pitcairn) lies well below the Samoa array.

The sample group on the high ¹⁴³Nd/¹⁴⁴Nd prong is also the lowest in ²⁰⁶Pb/²⁰⁴Pb and ²⁰⁷Pb/²⁰⁴Pb of all the shield lavas (Fig. 7). All Samoan lavas lie to the right (high ²⁰⁶Pb/²⁰⁴Pb side) of the terrestrial Pb Geochron and are in the mid-range of the elongate, worldwide OIB cluster; they are situated entirely above the Northern Hemisphere Reference Line (NHRL; Hart, 1984) in both ²⁰⁷Pb/²⁰⁴Pb and ²⁰⁸Pb/²⁰⁴Pb (Figs. 7 and 8). The most radiogenic ²⁰⁶Pb/²⁰⁴Pb compositions (19.4) are found not in the highest ⁸⁷Sr/⁸⁶Sr samples, but in samples from Vailulu'u Seamount (of moderate ⁸⁷Sr/⁸⁶Sr ~ 0.7055). On the other hand, the highest ²⁰⁷Pb/²⁰⁴Pb (15.65) and ²⁰⁸Pb/²⁰⁴Pb (39.8) correspond to the EM2-defining Malumalu lavas, implying that EM2 is an old reservoir of high time-integrated Th/U.

The Society array (not shown in Fig. 7) is much steeper, falling below the NHRL at low ²⁰⁶Pb/²⁰⁴Pb and crossing above it, to overlap the Malu trend data from Samoa. Interestingly, the highest ⁸⁷Sr/⁸⁶Sr sample from Tahaa (Societies) lies very close to our extreme ⁸⁷Sr/⁸⁶Sr

sample in ²⁰⁷Pb/²⁰⁴Pb - ²⁰⁶Pb/²⁰⁴Pb, but is far lower than it in ²⁰⁸Pb/²⁰⁴Pb. Note in Figure 8 that the Society array lies close to the NHRL, and is totally distinct from the Samoa field.

The ³He/⁴He ratios of Samoan lavas range from 8 R_A at high ⁸⁷Sr/⁸⁶Sr to a maximum of 26 R_A at generally lower ⁸⁷Sr/⁸⁶Sr (Fig. 9). New data support the existence a primitive helium mantle (i.e. PHEM of Farley et al., 1992) but with depleted Sr and Nd isotopic compositions (i.e. FOZO of Hart et al., 1992). With increasing ⁸⁷Sr/⁸⁶Sr, values of ³He/⁴He asymptotically approach ~8 R_A, showing that the helium isotopic composition of EM2 is approximately equivalent to that of MORB and much higher than the atmospheric values of recycled crustal materials (see discussion by Farley et al., 1992). This low ³He/⁴He value of EM2 is either inherent to the EM2 source, or is a product of diffusive equilibrium with the upper mantle over long timescales (see section 9).

The trace element character of the Samoan lavas display typical OIB features (Hofmann, 1988; Weaver, 1991), with trace element enrichments up to 100 times primitive upper mantle (PUM), the highest normalized concentrations at the highly incompatible elements, and negative anomalies at Cs, K and Pb (Fig. 10). The largest inter-volcano differences are at Pb, Rb, Ba, and Th. Weaver (1991) employed the trace element ratios Rb/Nb, Ba/Nb, Ba/Th. and Ba/La to distinguish between the EM1 and EM2 species. He used these trace elements to argue for a recycled sedimentary component as the cause for the EM signature, and ascribed the difference between EM1 and EM2 to a pelagic versus terrigenous sedimentary provenance. Therefore, a comparison between the Samoan lavas (extreme EM2) and those from Pitcairn (extreme EM1) should theoretically show the greatest differences in these ratios. However, recent studies on Pitcairn lavas (Dostal et al., 1998; Eisele et al., 2002) show nearly complete overlap with the Samoan lavas for Weaver's classification ratios, unlike the clear distinction between EM1 and EM2 previously reported for lavas with less extreme endmember signatures (Fig. 11). This result makes the trace element differences between EM1 and EM2 very difficult to resolve and discourages the description of EM1 and EM2 as having "pelagic" and "terrigenous" components, respectively. Furthermore, the Plank and Langmuir (1998) study of the compositions of sediment being subducted at today's convergent margins shows that pelagic and terrigenous sediments are: (1) not notably different in trace element ratios such as Ba/Th, and (2) not typically occurring alone in subducted sedimentary sections, but are instead components of the whole, mixed sedimentary package. Hence, generating mantle endmembers by recycling of only pelagic or only terrigenous sediment seems physically unlikely.

6.2 Shield vs. Post Erosional

As initially observed by Wright and White (1986), post-erosional (PE) lavas are isotopically distinct from all shield lavas. There is a commonality among the PE lavas from all along the chain (Savai'i, Upolu and Tutuila), in contrast to the observation that each island displays a unique isotopic birthmark in its shield lavas. The PE lavas show restricted ⁸⁷Sr/⁸⁶Sr values that plot mid-range in the Samoan field, have the lowest ²⁰⁶Pb/²⁰⁴Pb values and some of the lowest ²⁰⁸Pb/²⁰⁴Pb values of the whole sample suite (Figs. 6 and 8). The PE field on the ²⁰⁶Pb/²⁰⁴Pb versus ²⁰⁷Pb/²⁰⁴Pb plot (Fig. 7) is unusual, as it is elongate in an almost inverse direction to the shield trend (Wright and White, 1986).

Overall, the new Savaiian lavas are all of the same chemical nature as the post-erosionals, even though many are samples of the oldest-mapped flow series on the island (Fagaloa Series; Kear and Wood, 1959). These Savai'i lavas, as well as most other PE lavas, are clearly distinguishable from shield lavas by having the highest Nb/U and Ba/(La,Sm,Nb,Th) ratios of the whole sample suite (Fig. 12). Given the earlier discussion of the young radiometric ages for this "shield" series, we believe this sequence is in fact post-erosional, and not shield. The alternative explanation, that all of Savai'i is young and not part of an age-progressive Samoan hotspot track, is belied by the 2.05 my age for a trachyte cobble from the Vanu River valley (see above). Either way, we cannot rule out the possibility that PE lavas and shield lavas are geochemically the same on Savai'i, but nowhere else in Samoa.

What accounts for the distinct trace element and isotopic differences between shield and PE lavas? The commonality among Samoan PE lavas possibly derives from a similar history of being brewed and aged in the crust and lithosphere, unlike shield lavas that may have a shorter residence time in this shallow environment. Local Tongan sediments (from DSDP Site 595/596, about 1000 km southeast of Samoa) have Pb isotopic compositions (Plank and Langmuir, 1998) with the general characteristics of PE lavas (Figs. 7 and 8). Pb isotopic compositions of marine sediments are highly variable over short distances and other sediments could likely be found nearer to Samoa that provide closer fits to the Samoa post-erosional Pb field (which lies near the lower end of the general marine sediment array; Abouchami and Goldstein, 1995; O'Nions et al., 1998; Plank and Langmuir, 1998; Jones et al., 2000). In support of a sediment component in the PE lavas are values for δ¹⁸O of olivine (5.5-5.7‰; Eiler et al., 1997) which are elevated over upper mantle values and can be interpreted to reflect the heavy values documented for marine sediments (also see discussion

below). In other words, we cannot rule out the late-stage incorporation of modern marine sediments in PE lavas based solely on isotopic compositions. Trace element ratios may provide a stronger constraint on the presence or absence of a modern sediment component; one would expect the PE lavas to inherit the high Pb/Ce, high REE/HFSE, low Sm/Yb, and Ba-enriched ratios characteristic of both local and globally averaged marine sediments (see Fig. 17; Plank and Langmuir, 1998). This is not the case for the PEs, which have, of all suspected traits, only notably high Ba (Figs. 11 and 12).

6.3 Mixing Arrays

The spread of isotopic compositions in the Samoan lavas can be attributed to either (1) processes that generate an infinite number of chemical (i.e. parent/daughter) heterogeneities within the mantle that, upon long-term storage, evolve into an infinite number of isotopic heterogeneities or (2) processes that produce a small number of unique chemical compositions that, upon long-term storage, result in a limited number of "end-member" isotopic compositions available for mixing. In order for the first option to produce sub-linear arrays in 2-D and 3-D isotope space, there must be a single process which acts systematically to varying degrees or at various times. Hence, talk of or modeling of the most extreme values (i.e. endmember mantle components) is the same in either case.

The lavas from Malumalu undeniably establish the existence of a reservoir with high 87 Sr/ 86 Sr (at least 0.7089), low 143 Nd/ 144 Nd (at most 0.5125), and 206 Pb/ 204 Pb, 207 Pb/ 204 Pb, and ²⁰⁸Pb/²⁰⁴Pb values near 19.3, 15.65 and 39.9, respectively. An unaltered sediment reservoir can be immediately ruled out as the cause of the EM2 component in Samoan shield lavas: although Global Subducting Sediment (GLOSS; Plank and Langmuir, 1998) and local Tongan sediment (Site 595/596; Plank and Langmuir, 1998) each have convincing ⁸⁷Sr/⁸⁶Sr and ²⁰⁷Pb/²⁰⁴Pb compositions (Figs. 6 and 7), they are severely inadequate (low) in ²⁰⁶Pb/²⁰⁴Pb and ²⁰⁸Pb/²⁰⁴Pb to generate the isotopic signatures displayed by the shield lavas (Fig. 8). Therefore, for recycled sediment to have evolved to the EM2 coordinate in Sr-Nd-Pb isotope space, subduction zone alteration of ancient sedimentary packages needed to be very specific: U/Pb and Th/Pb must increase, while Rb/Sr and Sm/Nd remain very much the same. In the dehydration of subducted oceanic crust, this is shown to be the case for all systems except Rb/Sr: Rb is about 5 times more mobile than Sr (Ayers, 1998), so the final dehydrated product has significantly lowered Rb/Sr ratios. Experiments on the dehydration and melting of sediments (Johnson and Plank, 1999) give rather inconclusive results for relative trace element partitioning of these parent/daughter ratios, and suggest that partitioning can be extremely variable depending on the minerals present and the degree of dehydration.

Although the Samoan lavas are isotopically extreme, the "pure" EM2 signature may be even more extreme. For example, clinopyroxene and glass separates from peridotite xenoliths from Savai'i studied by Hauri et al. (1993) yield ⁸⁷Sr/⁸⁶Sr values up to 0.7128 and have been interpreted to represent metasomatism of oceanic lithosphere by a small degree carbonatitic melt (not diluted by mixing with depleted mantle) from the same source as that which provides melts for Samoan volcanism. However, the Pb isotopes in these rare xenoliths (²⁰⁶Pb/²⁰⁴Pb ~18.86; ²⁰⁸Pb/²⁰⁴Pb ~39.76) lie well outside the isotopic array set by the Samoan lavas (Fig. 8); this suggests an origin for the enriched component in these xenoliths from a smaller, unique reservoir, unrelated to extant Samoan lavas.

Clearly, though, EM2-rich samples are more rare than samples of a less-enriched nature. On a plot of ²⁰⁶Pb/²⁰⁴Pb against ⁸⁷Sr/⁸⁶Sr (Fig. 13), the Samoan samples can be enclosed in a triangle where the high ⁸⁷Sr/⁸⁶Sr apex is defined by EM2. At lower ⁸⁷Sr/⁸⁶Sr, there are two components, one with higher 206Pb/204Pb than EM2 and one with lower 206Pb/204Pb, but both assuredly depleted according to their high 143Nd/144Nd values (Fig. 6). The low ²⁰⁶Pb/²⁰⁴Pb, low ⁸⁷Sr/⁸⁶Sr apex (note the Upolu data cluster) has a signature tending toward DMM, but the strict use of the most depleted MORB/DMM isotopic values is not necessarily the only option for describing this component. The sub-Samoan upper mantle has been punctured by multiple mantle plumes in its 110 Myr lifespan, so may no longer be strictly, or homogeneously, pure DMM (see the South Pacific Isotopic and Thermal Anomaly; Staudigel et al., 1991). Also, we do not absolutely require the low 206Pb/204Pb depleted component to reside in the upper mantle (i.e. it could be part of the plume), although it's most easily visualized as being there given current notions of mantle dynamics. Regardless of these disclaimers, the use of anything but a generic DMM isotopic composition is arbitrary, and ultimately only compromises the generality of our observations and conclusions.

The high ²⁰⁶Pb/²⁰⁴Pb, low ⁸⁷Sr/⁸⁶Sr component (obvious in the Ta'u and Vailulu'u lavas; Fig. 13) is suggestive of mixing with a HIMU mantle component. This component may also be present in the Samoan plume, but there is reason to believe HIMU material has under-plated the Samoan lithosphere in the past. Calculated hotspot tracks show that 20-25 million years ago, the Cook-Austral plume was located beneath the lithosphere on which the Samoan Islands presently sit (Norton, 2000). The Cook-Austral chain shows great variation in isotopic compositions (Fig. 14), not all of which would fit the Samoan data in multi-

isotope space. However, there is one volcano, Raivavae, which has the isotopic compositions appropriate to be a significant component in the Vai Trend lavas (Fig. 14; data from GEOROC database); we are not suggesting that Raivavae itself is contributing to the Samoan lavas, but that isotopically similar material may be underplating the Samoan island chain.

A fourth mixing component must be acknowledged when considering ³He/⁴He values. Figure 9 shows the inverse relationship between ⁸⁷Sr/⁸⁶Sr and ³He/⁴He. The EM2 component can be classified as having a ³He/⁴He signal which asymptotically approaches the average DMM value of ~8 R_A (Kurz et al, 1982) at high ⁸⁷Sr/⁸⁶Sr. HIMU has also been shown to have low ³He/⁴He values (Graham et al., 1993; Hanyu and Kaneoka, 1997; Hilton et al., 2000) and likely explains why Vailulu'u (with the largest HIMU component) is in parallel with Malumalu on Figure 9. Therefore, all three endmember components discussed above have low ³He/⁴He, thus requiring an additional reservoir to account for high ³He/⁴He. High ³He/⁴He values are found in the center of the Samoan Sr-Pb data array, at Ta'u and Tutuila, and generally decrease towards the outer fringes (Fig. 13). Farley et al. (1992) named this component the primitive helium mantle (PHEM) but new data suggest this reservoir has depleted ⁸⁷Sr/⁸⁶Sr and ¹⁴³Nd/¹⁴⁴Nd (like FOZO of Hart et al., 1992), and not bulk-earth-like values assigned to PHEM.

All four mantle components are in the Samoan plume from a magmatic standpoint. But what material is coming from the deep mantle is another story. We can make a good case for the depleted component coming from entrainment of the widely documented depleted upper mantle and the radiogenic Pb component (HIMU-ish) coming from entrainment of under-plated lithosphere from the HIMU Cook-Austral chain. This means the deep mantle material within the Samoan plume is dominantly EM2 and PHEM/FOZO. The sequence of mixing these components is difficult to ascertain, as the length scale of compositional heterogeneity and differences in solidus temperatures (i.e. solid vs. melt mixing) are unknown.

6.4 Spatial/Temporal Evolution

Samoan shield samples on the ²⁰⁶Pb/²⁰⁴Pb - ²⁰⁸Pb/²⁰⁴Pb plot form two *en echelon* trends of positive slope (Fig. 8) which are most distinctly separated at high ²⁰⁶Pb/²⁰⁴Pb, and converge at lower ²⁰⁶Pb/²⁰⁴Pb. The isotopic trends correspond to the two topographic ridges of the Samoan islands (Fig. 2); for a given ²⁰⁶Pb/²⁰⁴Pb, the southern Malu Trend has higher ²⁰⁸Pb/²⁰⁴Pb than the northern Vai Trend. Within each of the two trends, isotopic enrichment

increases with decreasing age along the volcanic ridge. This relationship, shown clearly in a plot of distance versus ²⁰⁶Pb/²⁰⁴Pb (Fig. 15), has remarkable correlation and is striking in its implication of a systematic evolution of plume material or mantle processes. Figure 15 also shows how the Malu and Vai Trends form a continuum though time: even though each ridge independently displays isotopic enrichment with distance/time, the younger Vai Trend is generally higher in ²⁰⁶Pb/²⁰⁴Pb than the older Malu Trend (note that Malumalu may overlap in age with Ta'u and Vailulu'u). Of the four mixing components, low ²⁰⁶Pb/²⁰⁴Pb values are found only in the DMM reservoir (~18.0; Fig. 14). Therefore, the increase in ²⁰⁶Pb/²⁰⁴Pb with younging of volcanoes is interpreted to be a waning of the DMM component in the Samoan lavas, with a resulting increase in the abundance of EM2 and HIMU components. The separation of the Vai and Malu Trends in Pb-isotopic space indicates a higher HIMU/EM2 ratio in the Vai Trend.

Moving east along each of the two Trends, there are systematic increases in K/Na, Rb/Sr, La/Sm, La/Yb, Ba/Sm, Th/Nb, Th/Zr, Nb/Y, Nd/Sm, Nb/Zr, and U/Nb (Fig. 16); in other words, incompatible-element-enrichment increases with Pb isotopic enrichment, distance, and decreasing age. Due to correlations between isotopes and trace elements like those seen in Figure 16, variations in trace element ratios are easily attributed to differences in composition between the low ²⁰⁶Pb/²⁰⁴Pb source and the high ²⁰⁶Pb/²⁰⁴Pb sources. However, we are witness not to the source compositions, but to the products of "source processing". Because the process of melt generation has maintained (or not overly obscured) trace element correlations with isotopic compositions, we can infer some characteristics of the sub-Samoan mantle.

Possible explanations for the systematic chemical evolution of the Samoan plume include:

- 1. Horizontal zonation of plume material, implying a length-scale of heterogeneity on the order of volcano spacing, as has been suggested for the Hawaiian Islands (see below). In this case, trace element variations are truly source variations.
- 2. A lithologically-homogeneous mantle in which peridotite components of variable composition occur in the same proportions beneath all Samoa, but exist on a length-scale large enough to allow preservation of disequilibrium between the components. In this case, variable potential temperature of the plume would result in preferential sampling of components based on their respective solidus temperatures. Enriched materials would be sampled at small degrees of melting and trace element enrichment is partly a function of degree of melting.

 A vertically-stratified plume, changing composition and/or physical properties as upwelling proceeds, and affecting the degree of entrainment of ambient upper mantle and lithospheric assimilation.

In the Hawaiian Islands (an EM1 plume), isotopically distinct, topographic *en echelons*, named the "Kea" and "Loa" Trends, have also been documented (Tatsumoto, 1978; Staudigel et al., 1984; Abouchami et al., 2000). The Society Islands (another EM2 archipelago) display similar sub-parallel trends in both geographic and Pb isotopic space (using data compiled in the GEOROC database). However, nothing so temporally systematic as that in Samoa has been previously reported. Chemical zonation of a mantle plume (e.g. Kurz et al., 1995; Hauri et al., 1996; Lassiter et al., 1996; DePaolo et al., 2001) may explain isotopic lineaments within island chains, but fails to address how this chemical heterogeneity may translate into topographic features. On the other hand, creation of topographic lineation as a consequence of either 1) the lithosphere's structural response to loading (e.g. Hieronymus and Bercovici, 1999; Hieronymus and Bercovici, 2000) or 2) magma rising in "plumlets" instead of a continuous stream (Ihinger, 1995) ignores the fact of correlative chemical variations. Even so, some common dynamic feature clearly exists, independent of mantle taxonomy, for the way in which plumes forge through the mantle/crust, melt, and arrive at Earth's surface.

7. Calculation of a "Pure" EM2 Lava

The following calculation is aimed at defogging the trace element pattern for lavas of the enriched endmember, through "un-mixing" (subtracting) Ta'u lavas (average ⁸⁷Sr/⁸⁶Sr = 0.7046) from the most EM2-rich Malumalu lavas, under the assumption that the highest ⁸⁷Sr/⁸⁶Sr lavas are, instead of pure EM2 melts, still somewhat contaminated by melts from a depleted/less enriched mantle. As a group, Ta'u lavas are closest to the PHEM mixing component (Figs. 13 and 14). By this calculation, trace element differences between unenriched and enriched mantles are accentuated, and help to clarify the trace element characteristics of the EM2 source.

We extrapolate to the end-member trace element pattern of an EM2 melt in effect by subtracting the averaged trace element composition of Ta'u lavas from the Malumalu lavas until the ⁸⁷Sr/⁸⁶Sr composition equals 0.7128; this value derives from an analysis of cpx contained in a metasomatized peridotite xenolith from Savai'i (Hauri et al., 1993). Although

these xenoliths are not an extension of the Samoan Pb isotope array (Figs. 7 and 8), for lack of a better stopping point, they do place an upper limit on oceanic mantle Sr isotopic ratios.

Mixing between Ta'u and a "pure" EM2 component to make the most enriched Samoan samples (Malumalu samples 78-1 and 78-3) is calculated with the following two equations:

$${}^{(87}Sr/^{86}Sr)_{78-1} = \frac{F[Sr]_{EM2}(^{87}Sr/^{86}Sr)_{EM2} + (1-F)[Sr]_{Tau}(^{87}Sr/^{86}Sr)_{Tau}}{F[Sr]_{EM2} + (1-F)[Sr]_{Tau}}$$

$$(1)$$

$$[Sr]_{78-1} = F[Sr]_{EM2} + (1-F)[Sr]_{Tau}$$
(2)

The concentration of Sr ([Sr]) in EM2 and the fraction of the EM2 melt, F, are solved simultaneously so that the right hand of equation (1) equals the ⁸⁷Sr/⁸⁶Sr composition of the two extreme Malumalu lavas (0.70889). With the value for F, concentrations of all trace elements can be calculated for the EM2 melt by using the structure of equation (2) and are reported in Table 7. Lava compositions used in this calculation have been corrected for crystal fractionation by incremental addition of olivine (or subtraction in the case of 78-1, 78-3 and 74-1) until the melt compositions reaches a Mg# of 73 (olivine addition ranges from 10-51%; olivine subtractions are 10%, 7% and 23%, respectively). Note from Figure 4 that Ta'u and Malumalu have very similar crystal fractionation trajectories with minimal cpx loss. All Ta'u samples have been utilized except for T21 which is plagioclase-rich and T44 which is an ankaramite.

The resulting fraction of EM2 "melt" in the Malumalu "mixture" is 51%, and the resulting ¹⁴³Nd/¹⁴⁴Nd ratio for the EM2 component equals 0.51235. Figure 17 shows the trace element pattern for the calculated "pure" EM2 melt component; note enrichments at Rb and Th that are almost 120 times PUM, negative anomalies at Cs and Ba, and an almost non-existent Pb anomaly. The REE slope of the calculated EM2 melt is steeper than both Malumalu and Vailulu'u, and the overall trace element pattern from U to the right is remarkably smoother than either the Malumalu or Vailulu'u pattern, save for dips at Sr and Ti. In general, the degree of enrichment in the EM2 melt is greatest for the highly incompatible elements.

The calculated trace element pattern of the "pure" EM2 melt is compared to: 1) an estimate of global subducting sediment (GLOSS; Plank and Langmuir, 1998) and 2) a local

sediment from DSDP Hole 595/596 analyzed for the GLOSS compilation (Fig. 17). Clearly, the sediment trace element patterns are very different from the calculated EM2 component. In particular, the sediment spidergrams are marked by large negative anomalies of the high-field-strength elements (HFSE; Nb, Ta, Zr, and Hf), and large positive Pb and Ba anomalies, whereas the calculated Samoan enriched component has no such features; in fact, the Ba anomaly becomes more *negative* in the EM2 melt. Also, the heavy rare-earth-element slope of the EM2 melt is significantly steeper than the sediment patterns: Sm/Yb for the sediments is 2.1 whereas for the EM2 melt is 7.2. The only argument in favor of sediment addition is the significantly decreased Pb anomaly in the EM2 melt. However, we (1) do not believe this alone lends credence to the sediment theory, and (2) show in our non-sediment model below how Pb in the EM2 source does not have a negative anomaly.

Ultimately, the calculated EM2 spidergram is inconsistent with standard models invoking ancient sediment recycling to explain the enrichment of the EM2 mantle source. As discussed below, it is unlikely that any chemical processing during subduction would so effectively "smooth out" the typically jagged spidergram of oceanic sediment. Alternatively, if the enriched plume material is argued to derive from addition of present-day sediments, the trace element patterns of local sediments should be directly reflected in the EM2 melt and they are not. Therefore, late-stage contamination of plume material with local sediment is also an unsatisfactory explanation for the observed chemical characteristics of the enriched Samoan basalts (and this point is strongly supported by the Pb isotope evidence shown in Figs. 7 & 8). Production of the EM component by deep mantle fractionations involving high-pressure phases such as Ca or Mg perovskite likewise will lead to jagged, not smooth, spidergrams (Hirose et al., 2004). Segregation of carbonatitic melts from mantle assemblages has been used to explain elevated trace element concentrations in oceanic lavas (see Zindler and Hart, 1986), but this process also causes irregular trace element patterns (e.g. Klemme et al., 1995; Sweeney et al., 1995; Hoernle et al., 2002). Instead, the remarkably smooth EM2 melt spidergram gives the uncanny impression of having originated from nothing but "unadulterated" melting processes within the upper mantle.

8. Sediment Recycling?

Osmium and oxygen isotopes are thought to be "smoking guns" for sediment/slab recycling (Eiler et al., 1997; Shirey and Walker, 1998; van Keken et al., 2002). Due to the

incompatibility of Re (Righter and Hauri, 1998) and compatibility of Os (Hart and Ravizza, 1996) in mantle melting, elevated Re/Os ratios in crustal materials should evolve to radiogenic osmium during long-term storage within the mantle. Altered upper MORB crust and marine sediments are enriched in heavy oxygen (δ^{18} O of ~15-25%; Savin and Epstein, 1970; Lawrence et al., 1979; Staudigel et al., 1995; Alt, 2003) by low-temperature fractionation processes at the Earth's surface. This is high above the δ^{18} O value of 5.2% for upper mantle olivine (Ito et al, 1987; Mattey et al., 1994; Eiler et al., 1997). Therefore, the standard theory for the origin of EM2 involving recycling of mafic crust plus terrigenous sediment would suppose Samoan lavas to have both elevated δ^{18} O and δ^{18} Os compositions.

Eiler et al. (1997) demonstrated that EM2 basalts from Samoa (Savai'i post-erosional) and the Societies do have the highest δ^{18} O of all OIB's (δ^{18} O of olivine up to 6.1‰), explainable by the incorporation of ~5% terrigenous sediment addition to DMM. Using values chosen by Eiler et al. (1997) for the concentrations of Sr, Nd, and Pb in DMM and sediments, the sediment contribution to the trace element budget in the EM2 source will be 50%, 68% and 96%, respectively, for these elements. Clearly then, the trace element pattern of EM2 lavas should reflect the trace element patterns of sediment, but they do not (see Fig. 17). Eiler et al. (1997) also mention the possibility that metasomatism can elevate δ^{18} O values in magmas, and the present work recommends this idea be further explored.

Osmium isotopic compositions are likewise not so "smoking" of a sediment component. Combining data presented here (Table 4) with those from Hauri and Hart (1993), Samoan basalt samples with >80 ppt Os (ranging in ⁸⁷Sr/⁸⁶Sr from 0.7046 to 0.7089) reveal ¹⁸⁷Os/¹⁸⁸Os ratios of 0.124 - 0.130 which do not correlate with any other isotope system. Samples with <80 ppt Os (5 out of 21 in total) have elevated ¹⁸⁷Os/¹⁸⁸Os ratios and are interpreted to be contaminated with seawater (see Shirey and Walker, 1998). The small range in ¹⁸⁷Os/¹⁸⁸Os compositions of pristine samples spans values estimated for the primitive upper mantle (0.129; Meisel et al., 1996) and DMM (~0.125; Standish et al., 2002), and is much lower than the upper limit of 0.16 displayed in HIMU and EM1 lavas (Hauri and Hart, 1993; Reisberg et al., 1993; Eisele et al., 2002).

The unradiogenic ¹⁸⁷Os/¹⁸⁸Os values for these Samoan lavas represent either (1) a similarly unradiogenic mantle source, or (2) re-equilibration of more radiogenic Os components with unradiogenic upper/lower mantle through special processes that are not active beneath HIMU or EM1 hotspots. With regard to the former option, and to test the standard model, low Os concentrations in sediments may prevent a sediment component

from significantly elevating ¹⁸⁷Os/¹⁸⁸Os ratios in the EM2 source. In a simple case, if DMM with ¹⁸⁷Os/¹⁸⁸Os = 0.125 and [Os] = 3000 ppt is mixed with sediment having ¹⁸⁷Os/¹⁸⁸Os = 1.0 and [Os] = 30 ppt (Peucker-Ehrenbrink and Jahn, 2001), then 35% of sediment is needed to change ¹⁸⁷Os/¹⁸⁸Os from 0.125 to 0.130. Here we are again left with an EM2 source whose trace element budget would be dominated by sediment, but do not observe such trace element patterns in the EM2 lavas nor see the implied correlations with other isotope systems. The second option, suggesting the Os budget derives from reequilibration, can be ruled out since olivine phenocrysts are in approximate equilibrium with coexisting liquids (Jackson et al., unpubl.) and have high ³He/⁴He ratios (i.e. are not xenocrystic, but rather truly phenocrystic). We conclude that the mantle sources for Samoan lavas all have inherently unradiogenic ¹⁸⁷Os/¹⁸⁸Os values and are not influenced by a sediment/crustal component.

Although slab/sediment recycling has been a common theory for the origin of EM2 for over two decades (see Introduction), there are major flaws in this train of thought. The Standard Model for generating the EM2 reservoir involves the introduction into the deep mantle of 1) oceanic crust which has been depleted of fluid-mobile elements, such as the large-ion-lithophile elements (LILE; e.g. Cs, Rb, K and Pb), by dehydration and 2) a relatively pristine (i.e. elementally unfractionated) continental crust component (i.e. terrigenous sediments). Although not typically considered in the Standard Model, it seems logical that trace elements of subducted sediments (pelagic and/or terrigenous) must be fractionated by the same process by which the subducted ocean crust is fractionated (dehydration) - especially since sediments are closer to the mantle wedge and likely to have greater water contents than the altered ocean crust. Whereas there have been experimental studies showing high trace element mobility during dehydration of subducted ocean crust (especially for the isotopically important elements Rb and Pb; see Ayers [1998] and Stracke et al. [2003] for overviews), very little similar work has been done on dehydration of subducted sediments (i.e. Johnson and Plank, 1999). Actually, there is growing geochemical evidence that not only a fluid component, but also partial melts of subducted sediments contribute to arc magmas. The high recycling efficiencies (up to 40%) of elements which are not particularly fluid mobile, such as Be, Th and Nd (see discussion by Johnson and Plank, 1999), suggest sediment melting is a reality, even though many thermal models predict subsolidus temperatures within the subducted sediment column (e.g. Peacock, 1996). Regardless of the mechanism of trace element fractionations in subducted sediments, it is clear that fractionations will occur and will result in significant loss of incompatible

elements, and a decrease in the mass of a possible future EM2 reservoir. Ultimately, it is grossly inconsistent to use modern, surface sediment as an approximation of the trace element and isotopic composition of a "sediment" component in the mantle – once subducted, the sediment will never look the same, especially for parent/daughter ratios like Rb/Sr and Th/Pb.

Additionally, since today's surface, terrigeneous sediments represent what has been extracted by convergent margin volcanism and/or continental crust formation, it is the residue, or complement, to surface sediments which should be our concern for what material is actually recycled deep into the mantle. For example, depletion of the fluid immobile elements Na and Ta in arc volcanics (Pearce and Peate, 1995), and hence sediments (Plank and Langmuir, 1998), will be matched by Nb-Ta enrichments in the material that is ultimately introduced to the deep mantle. Experiments on partitioning between dehydration fluids and eclogite mineral assemblages (garnet, clinopyroxene and rutile) suggest that depletion of high field-strength elements (including Nb-Ta) in arc volcanics is due to their high compatibility in residual rutile (Stalder et al., 1998) and is therefore not a sediment signature. Enrichment of HFSE in the subducted slab will offset HFSE depletions in the subducted sediment. This is why decreasing Nb anomalies with increasing ⁸⁷Sr/⁸⁶Sr ratios, as documented for EM1 and EM2 lavas by Eisele et al. (2002), are not supporting evidence for sediment recycling.

We believe there is an alternative explanation for correlation between Nb anomalies and isotopic compositions. Figure 18 shows Nb/Nb* (calculated as Nb_N/√(Th_N×La_N); Eisele et al., 2002) plotted with ²⁰⁸Pb/²⁰⁴Pb and La/Sm ratios of lavas from Samoa and Pitcairn. We have used ²⁰⁸Pb/²⁰⁴Pb as a measure of EM2 abundance instead of ⁸⁷Sr/⁸⁶Sr only because it provides better correlations. Samoan lavas show inverse relationships between Nb/Nb* and ²⁰⁸Pb/²⁰⁴Pb as well as La/Sm. Pitcairn lavas (from Eisele et al., 2003) show a negative correlation between Nb/Nb* and La/Sm, which overlaps with the Samoan lavas, and a more shallow slope than Samoa for Nb/Nb* against ²⁰⁸Pb/²⁰⁴Pb (the greatest isotopic variation in the Pitcairn lavas is in ¹⁴³Nd/¹⁴⁴Nd). Pitcairn and Samoa samples have almost an identical range in both La/Sm and Nb/Nb*, even though the isotopic variability is greater in Samoa. Also plotted in Figure 18 is a trajectory for variable degree of melting of a depleted mantle, showing that small changes in F can produce large changes in both La/Sm and Nb/Nb*. Therefore, variable Nb/Nb* (previously interpreted as only a source effect) can be produced by recent variations in melt production, and is most likely what causes (1) scatter in the plots of Figure 18 and (2) the same Nb/Nb* variation in Pitcairn as Samoa given less isotopic

variation. The correlation of ²⁰⁸Pb/²⁰⁴Pb (and ⁸⁷Sr/⁸⁶Sr) with Nb/Nb* can be interpreted as an ancient enrichment of mantle by a small degree (low Nb/Nb*) melt, as suggested by the calculated EM2 melt and modeled below.

9. Metasomatic Origin of EM2

Given the many failures of the "sediment recycling" model for EM2, as enumerated above, we propose here a new model that invokes metasomatic enrichment of ancient oceanic lithosphere, followed by long-term storage in the deep mantle and recent return to the surface as the Samoa plume. Conceptually, this model derives from the autometasomatic process proposed by Zindler et al. (1979) and Roden et al. (1984). Numerous authors have appealed to metasomatism of oceanic plates to generate chemical heterogeneities that can be tapped prior to plate subduction (Hawkesworth et al., 1979, 1984; Halliday et al., 1992; Class and Goldstein, 1997; Niu et al., 1996). Recycling of such metasomatized lithosphere, after long-term storage in the mantle, has been advocated by Richardson et al. (1982) and Niu and O'Hara (2003) as a source for enriched OIB.

If we start with the assumption that EM2 is a two-stage differentiate of bulk-earth, the slope on the ²⁰⁶Pb/²⁰⁴Pb - ²⁰⁷Pb/²⁰⁴Pb plot (Fig. 7) yields an age of 2.5 Ga. This is an age older than the commonly quoted average mantle differentiation age of 1.8 Byr (Hart, 1984). At that time, the composition of the mantle would have been more similar to primitive upper mantle than to the depleted mantle observed today (i.e. DMM). Assuming plate tectonics was operating 2.5 billion years ago in much the same way as it is today, this more primitive mantle material would have undergone depletion by melt extraction during upwelling under spreading ridges, then "turned the corner" and solidified to become depleted lithospheric mantle.

In the following calculations, we model the case in which small degree, deep melts not extracted at the ridge crest percolate up through the asthenosphere and impregnate the overlying lithosphere that had just undergone melt extraction on the ridge crest. This is essentially a metasomatic process. This metasomatized lithosphere then is recycled and stored in the mantle to become today's EM2 reservoir (Fig. 19). The melt fraction, amount of melt impregnation, and ratio of garnet to spinel peridotite melting are calculated so as to match parent/daughter ratios of EM2 for the Rb-Sr, Sm-Nd, U-Pb, Th-Pb and Lu-Hf systems, based on evolution from bulk earth 2.5 billion years ago. Bulk partition coefficients used for melting a primitive mantle source (McDonough and Sun, 1995) are

based on a compilation of D's from Kelemen et al. (2003) for melting of garnet and spinel peridotite, with the few exceptions listed in Figure 20. Bulk partition coefficients are weighted 72% garnet peridotite to 28% spinel peridotite. The best match to parent/daughter ratios is with a 1.1% impregnation of a depleted lithosphere by a 0.5% batch melt of a primitive mantle. The lithosphere represents a mantle depleted by 2% melt extraction, as calculated using the method of Workman and Hart (2003) and as reported in Table 8. Figure 20 shows the resulting trace element pattern of the EM2 source (also see Table 8). Rb/Sr has been fit to within <1%, Sm/Nd and Lu/Hf have been fit to within 3%, and Th/U has been fit to within 4%. The "unfortunate fits" are for U/Pb and Th/Pb, which are 53% and 58% too high respectively in the calculated EM2 source. This is clearly more a Pb problem than anything else. If the compatibility of Pb is lower by about a factor of two, as suggested by experimental partitioning data (Hauri et al., 1994; Salters et al., 2002), the U/Pb and Th/Pb ratios may be more precisely modeled. Because the mass fraction of melt added to the FOZO lithosphere (1.1%) is twice the degree of melting (0.5%) required to generate that impregnating melt, the mass of the mantle which melts must be twice as large as the mass of the metasomatized lithosphere.

Does this source lead to the observed ${}^{3}\text{He}/{}^{4}\text{He}$ values of 8 R_A for EM2? Given the general trace element enrichment in the impregnating melt, and making the standard assumption of extreme incompatibility of He, it is likely that the calculated EM2 source would have high He/U ratios and hence evolve to ${}^{3}\text{He}/{}^{4}\text{He}$ values higher than 8 R_A. There are two possible solutions. One concerns the relative compatibility of He and U; if at high pressure and low degree of melting He is more compatible than U (this has not been proven nor disproven), then the impregnating melt will have low He/U and potentially evolve to DMM-like ${}^{3}\text{He}/{}^{4}\text{He}$ values (by coincidence). The second option is that the EM2 "package" has had a residence time in the upper mantle long enough (~1-2 Ga) to result in diffusive equilibrium of He (see model by Moreira and Kurz, 2001, for example); this option has obvious implications for the primary home of recycled lithosphere.

Although the above model leaves several questions unanswered, such as the scale-length of the heterogeneities created by the metasomatism, and the resulting lithologies (mafic veins or enriched peridotite), it is successful in producing the observed isotopic and trace element characteristics of the Samoan mantle source. It does not require *ad hoc* chemical manipulations in the subduction zone, as does the standard crust/sediment-recycling model. In fact, as the enrichment zone is limited to the lower parts of the lithosphere, it will be nearly invulnerable to subduction zone processing. It calls on a process for which there is

abundant evidence, particularly in the sub-continental lithosphere (Frey and Green, 1974; Menzies and Murthy, 1980; Menzies, 1983; Menzies and Hawkesworth, 1987). And insofar as small-degree melts are ubiquitous in the upper oceanic asthenosphere, the process is virtually guaranteed. We note also that the small-scale convection usually invoked for this part of the mantle (i.e. Richter rolls) provides an efficient means of upward advection of standing melt fractions, as well as the consequent decompression that will augment the melt fractions and facilitate melt/solid segregation.

10. Conclusions

A large suite of recently collected basalts from the Samoa hotspot chain have been analyzed for Sr, Nd, Pb, Os and He isotopes, and major and trace elements. Localities include the sub-aerial islands of Savai'i, Upolu and Ta'u, and the submarine seamounts Muli, Malumalu and Vailulu'u.

- 1. Samoan basalts are isotopically (Sr-Nd-Pb) the most extreme EM2 lavas in the oceanic database (⁸⁷Sr/⁸⁶Sr up to 0.7089). The Samoan isotopic arrays can be explained by mixing among four mantle components: DMM, EM2, HIMU and PHEM/FOZO. The deep plume material is most likely composed of EM2 and PHEM/FOZO, whereas the HIMU and DMM components are entrained into the plume in the upper mantle.
- 2. Systematic temporal and spatial variations in lava chemistry occur while going from west (older) to east (younger) along the chain: e.g., ²⁰⁶Pb/²⁰⁴Pb, La/Sm, Rb/Sr, Th/Zr. This indicates a waning of the DMM component and waxing of the EM2 and HIMU components in Samoan volcanoes over the last few million years.
- 3. The standard recycling model (ocean crust plus terrigenous sediment) fails as an explanation for producing Samoan EM2, as witnessed by the smooth spidergrams for EM2 lavas with negative Ba anomalies, low ¹⁸⁷Os/¹⁸⁸Os ratios, high ³He/⁴He (>8Ra) and mismatched Pb isotopic compositions.
- 4. The EM2 mantle source can be successfully modeled with the ancient (2.5 Ga) formation of metasomatised oceanic lithosphere, followed by storage in the deep mantle and return to the surface in the Samoan plume.

Appendix 2-1

Samples analyzed by M. Regelous were crushed in a steel jaw crusher to 3-5 mm sized chips, washed in deionised water, dried and handpicked in order to avoid chips which were visibly altered, or which contained vesicles. A portion of these chips was set aside for isotope analysis, the rest was powdered in an agate swing mill. Major element analyses were carried out by X-ray fluoresence at the Universität Mainz, Germany, using a Phillips PW 1404 instrument. Trace element concentrations were determined by ICPMS using a Fisons Plasmaquad II instrument at the University of Queensland, Australia. Full details of the procedure are given in Niu and Batiza (1997). The external precision on the concentrations of most of the trace elements measured is < 3%. The long-term average values for the BHVO-1 rock standard are reported in Table 5. Pb isotope measurements were carried out at the Max Planck Institut für Chemie, Mainz, following the procedure outlined by Abouchami et al. (2000). Between 50 and 100 mg of rock chips were washed in deionised water in an ultrasonic bath, then ultrasonicated in 6M HCl for 15 minutes, before being leached in hot 6M HCl for 1 hour. The HCl leachate was pipetted off, and the residue was rinsed, soaked in deionised water for 15 minutes, rinsed again and dried. This leaching procedure appears to remove much of the non-magmatic Pb that is contained in lessresistant components (surface contamination or alteration products), as discussed by Abouchami et al. (2000) and Eisele et al. (2003). The leached residues were dissolved in HF-HNO3, treated repeatedly with HNO3 and HCl until completely in solution, and Pb separated on anion exchange resin using HBr-HNO₃ mixtures (Abouchami et al., 2000). All reagents used were double-distilled, and total precedural blanks for the Pb chemistry were below 50pg. The eluent from the Pb columns was twice evaporated to dryness with 15M HNO₃, and redissolved in 3M HNO₃. Sr and Nd were separated from this fraction at the University of Bristol, U.K., using methods adapted from Pin et al. (1994). The sample in 3M HNO₃ was loaded onto columns containing 0.15ml of TRU spec resin, positioned so as to drip directly into a second column containing 0.1ml of Sr spec resin. After rinsing with 3M HNO₃, the columns were separated, and Sr was eluted from the Sr spec column in H₂O. The light- and middle-rare earth elements were recovered from the TRU spec resin by rinsing with 2.5M HCl. Nd was separated from this fraction using conventional HDEHP columns and 0.3M HCl. Pb isotope analyses were carried out using a triple spike technique to correct for instrumental mass fractionation. About 5% of the purified Pb fraction was transferred to a second beaker and spiked with a 204Pb-206Pb-207Pb triple spike. The spiked and unspiked fractions were loaded onto separate Re filaments with silica gel-H₃PO₄. Isotope compositions were measured using a Finnigan MAT-261 multicollector mass spectrometer (M.P.I. Mainz) in static mode, and the data for spiked and unspiked fractions were combined off-line to obtain the fractionation-corrected Pb isotope composition of the sample (Galer, 1999). During this study, the NBS981 Pb standard gave ²⁰⁶Pb/²⁰⁴Pb, ²⁰⁸Pb/²⁰⁴Pb ratios of 16.9403±0.0022, 15.4974±0.0020 and 36.7246±0.0058 respectively (2s, n=19). Sr and Nd isotope measurements were carried out on a Finnigan Triton multicollector mass spectrometer (University of Bristol) in static mode, and within-run exponential fractionation corrections applied using 86 Sr/ 88 Sr=0.1194 and 146 Nd/ 144 Nd=0.7219. The NBS987 Sr and J&M Nd standards gave 87 Sr/ 86 Sr and 143 Nd/ 144 Nd ratios of 0.710247±0.000008 (2 sigma, n=15) and 0.511113±0.000004 (n=12) respectively, during the period of the sample measurements.

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Figures

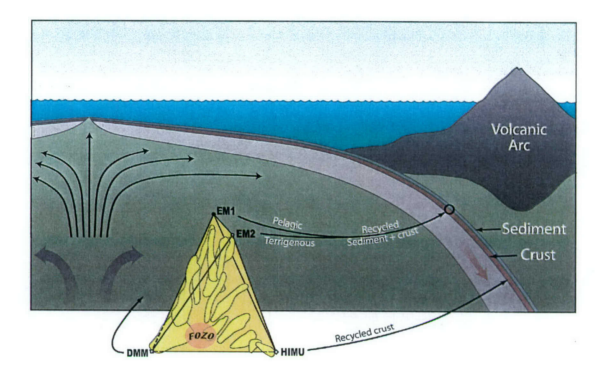


Figure 2-1. Schematic diagram of the Standard Model for the origin of isotopically defined mantle components. DMM (the Depleted MORB Mantle) is the melt-depleted upper mantle that supplies melts to mid-ocean ridges; HIMU (high U/Pb mantle) is a reservoir derived from recycling and long-term storage (billions of years) of oceanic crust; EM1 and EM2 are derived from recycling and long-term storage of oceanic crust along with pelagic or terrigenous sediment, respectively. Major contributions to the model have been from Armstrong (1968), Chase (1981), Hofmann and White (1982), Cohen and O'Nions (1982), White (1985), Zindler and Hart (1986), Weaver (1991), and Hart et al. (1992).

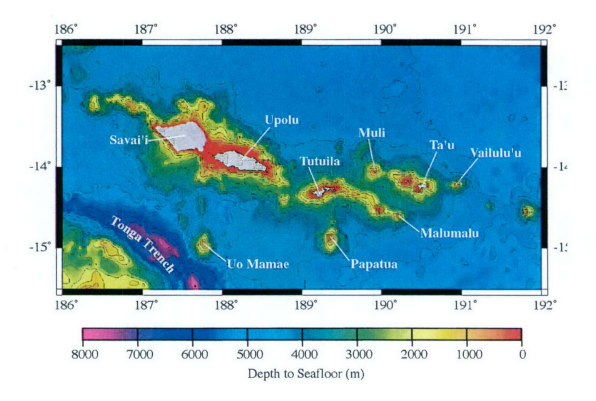


Figure 2-2. Bathymetric map of the Samoan volcanic chain made from merging inferred bathymetry from Smith and Sandwell (1994) with ship-track data from both the AVON 2/3 cruise (see Hart et al., 2000) and the GEODAS track-line database. Western Samoa is comprised of the two western islands, Savai'i and Upolu; American Samoa is comprised on Tutuila, Ofu, Olosega, and Ta'u. In the southwest corner of the map, where depths are down to 8000 m, is the northern termination of the Tonga Trench. Just off to the west at about 14.5°S is a transform fault bounding the Lau Backarc Basin to the south.

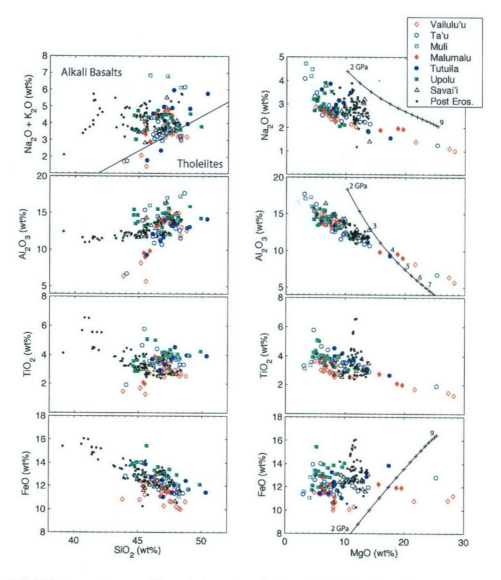


Figure 2-3. Major element compositions of Samoan basalts. Plots include data from Hauri and Hart (1997) for Savai'i lavas. Alkali-tholeiite line is from MacDonald and Katsura (1964). Trajectories of compositions for primary melts from fertile peridotites are plotted on some of the MgO diagrams, using the algorithms of Herzberg and Zhang (1996) in the pressure range of 2-8 GPa (tick marks every 0.5 GPa).

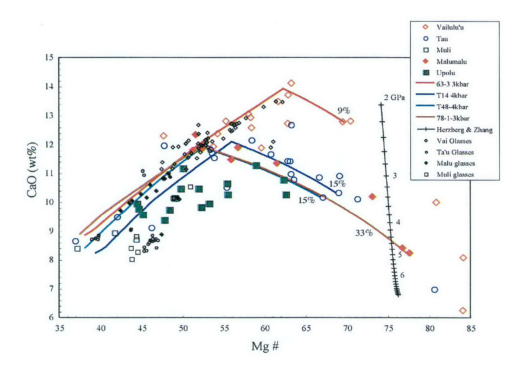


Figure 2-4. CaO plotted with Mg#'s for Samoan lavas. Mg# is calculated as molar percentage of Mg/(Mg+Fe²⁺) where Fe²⁺ is taken to be 85% of reported FeO. Compositions of primary melts from fertile peridotite are plotted using algorithms from Herzberg and Zhang (1996) in the pressure range of 2-8 GPa; tic marks are every 0.2 GPa. Crystal fractionation trends have been calculated using pMELTS at pressures of 3 and 4 kbar for best fits to compositional trends starting with some of the most MgO-rich lavas. Mass of olivine crystallized (expressed as a percent of the total initial mass) before clinopyroxene saturation is noted at the high Mg# end of the liquid lines of descent. Primary melts can be interpreted to have integrated depths of melting from 2.5-6 GPa, but CaO variations in the lavas more likely represent CaO contents of a heterogeneous mantle source. Plot includes data from Hauri and Hart (1997) for Savai'i lavas.

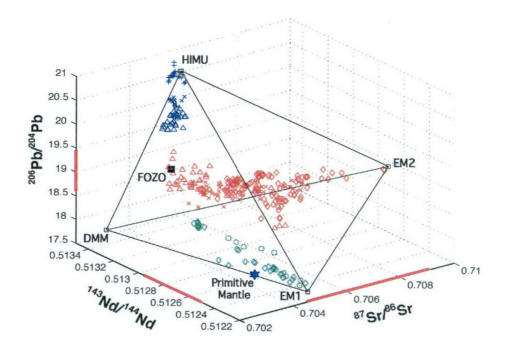


Figure 2-5. Mantle tetrahedron of Hart et al. (1992). Arrays from endmember defining island chains have been plotted using the GEOROC database and data presented in this manuscript. Island chains plotted for HIMU are in blue and include Tubuaii (×), Mangaia (+) and St. Helena (\triangle). EM1 islands are in green and include Pitcairn (\Diamond) and Walvis Ridge (\bigcirc). EM2 islands are in red and include Samoa (\Diamond), Societies (×), and the Marquesas (\triangle). Red bars along the axes mark the range of values for the Samoan Islands. EM2 has been extended from its previous coordinate (Zindler and Hart, 1986) to values for ${}^{87}\text{Sr}/{}^{86}\text{Sr}$, ${}^{143}\text{Nd}/{}^{144}\text{Nd}$, and ${}^{206}\text{Pb}/{}^{204}\text{Pb}$ at 0.7090, 0.5125, and 19.3, respectively.

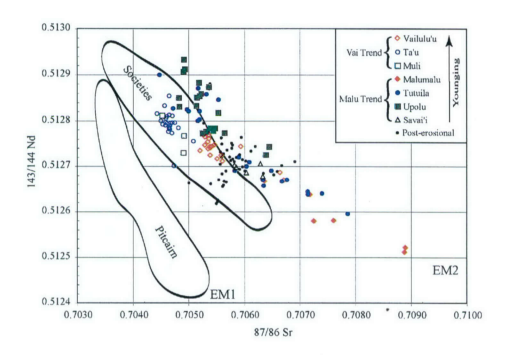


Figure 2-6. Sr and Nd isotopes for Samoan lavas. This as well as other isotope plots includes data from Wright and White (1987), Farley et al. (1992), and Hauri and Hart (1993). The legend here applies to all other isotope plots. The Vai Trend and Malu Trend correspond to topographic ridges of the volcanic chain (see Fig. 2). Savai'i samples marked with triangles are all from the Fagaloa Volcanic series. Post-erosional lavas include samples from Upolu and Savai'i. Fields for the Societies and Pitcairn were obtained from the GEOROC database. Coordinates for Globally Subducting Sediment (GLOSS) and local Tongan sediment are from Plank and Langmuir (1998).

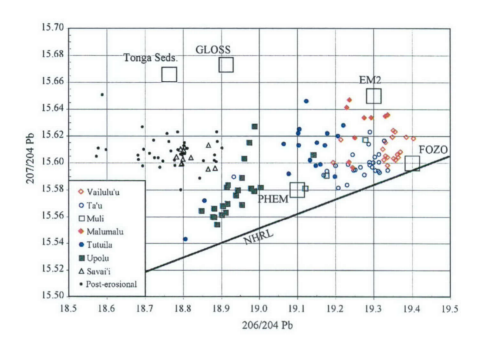


Figure 2-7. Plot of ²⁰⁶Pb/²⁰⁴Pb with ²⁰⁷Pb/²⁰⁴Pb of Samoan lavas. The Northern Hemisphere Reference Line (NHRL) lies significantly below the EM2 coordinate. Here, the Vai and Malu topographic lineaments can be distinguished as separate isotopic trends. Note how the post-erosional lavas are askew to the overall array of shield lavas. GLOSS = Globally Subducting Sediment (Plank and Langmuir, 1998); PHEM = Primtive Helium Mantle (Farley et al., 1992). Hauri et al. (1993) xenolith data derives from cpx and glass separates from Savaiian xenoliths. See Figure 6 for other references.

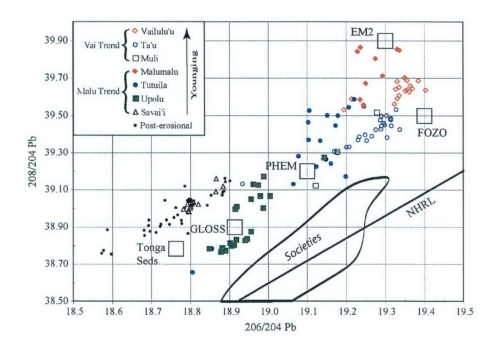


Figure 2-8. Plot of $^{206}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$ of Samoan lavas. Again, the Vai and Malu Trends are separated into two isotopic arrays. Along each trend, the age of volcanoes increases in the direction of lower $^{206}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$. See Figure 6 for references.

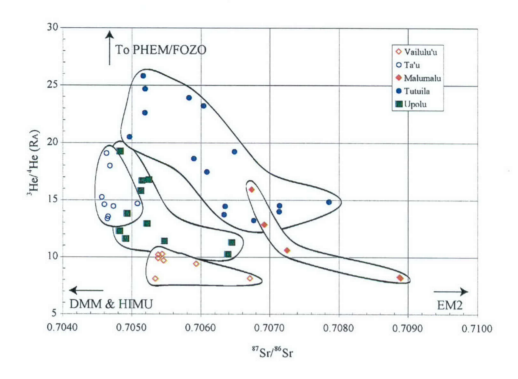


Figure 2-9. Plot of 87 Sr/ 86 Sr compositions of Samoan basalts with 3 He/ 4 He (R_A) of olivine phenocrysts and submarine glasses obtained from the same basalts. Some Tutuila samples are from Farley et al. (1992). EM2 is shown here to approach the DMM 3 He/ 4 He value of \sim 8 R_A at high 87 Sr/ 86 Sr.

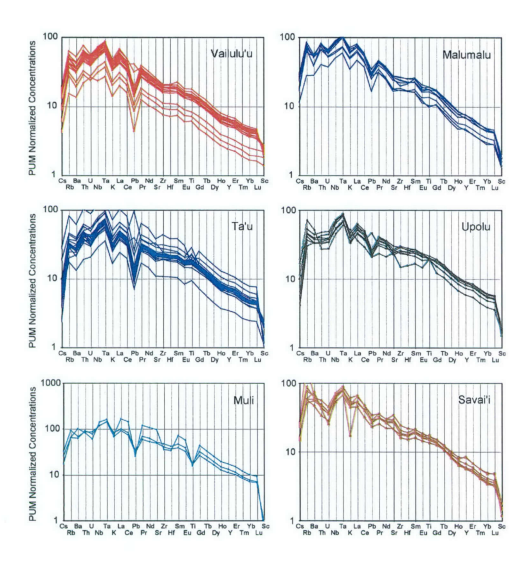


Figure 2-10. Trace element concentrations of Samoan lavas normalized to primitive upper mantle (PUM) of McDonough and Sun (1995). Note the difference in scale for the Muli lavas. Low concentration patterns are typically picrites (for example, the lowest three samples from Vailulu'u and lowest one from Ta'u). The highest concentration sample from Ta'u is T21, with 50% plagioclase phenocrysts.

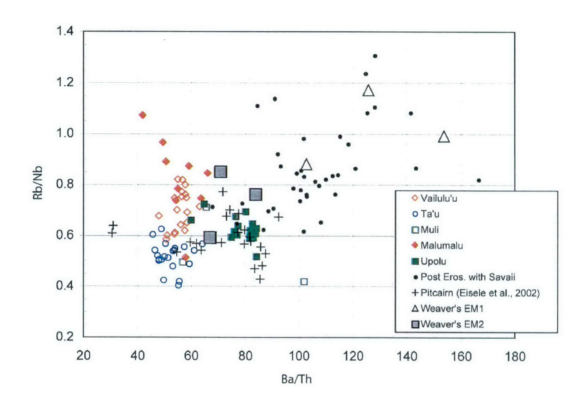


Figure 2-11. Ba/Th vs. Rb/Nb for lavas from Samoa (this study; Regelous et al., unpubl.; Hauri and Hart, 1997) and Pitcairn (Eisele et al., 2003) showing that Weaver's (1991) distinction between EM1 and EM2 trace element characteristics do not hold up to comparisons of lavas from endmember defining island chains (see Fig. 5). Pitcairn and Samoa show complete overlap in Ba/Th and Rb/Nb, whereas Weaver (1991) showed separate fields for EM1 and EM2 lavas. Plank and Langmuir (1998) report that terrigenous and pelagic sediments have indistinguishable Ba/Th ratios, each with a range of 10-220, with exceptions being rare hydrothermal clays and hemipelagic clays that are heavily-enriched in Ba. Therefore, the reason for initially identifying EM1 and EM2 as having recycled "pelagic" and "terrigenous" sediment, respectively, proves unfounded with further data collection.

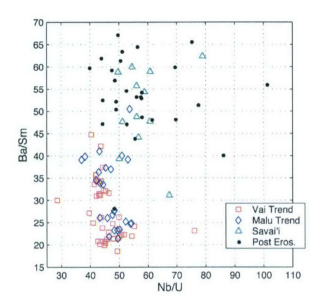


Figure 2-12. Plot of Nb/U vs. Ba/Sm used to highlight the trace element differences between shield and post-erosional lavas in Samoa. The new Savai'i lavas, sampled from the oldest mapped volcanic series on the island (Fagaloa Series; Kear and Wood, 1959), plot in the same field as post erosional lavas from all along the Samoan chain. This leads to the conclusion that either post-erosional lavas and shield lavas are the same on Savai'i, or post-erosional volcanism has been unusually extensive.

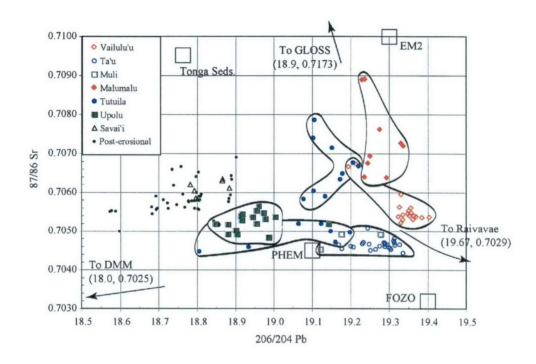


Figure 2-13. Sr and Pb isotope plot showing two classes of volcanoes – those which are elongate on the ²⁰⁶Pb/²⁰⁴Pb axis (Upolu, Tutuila Pago shield, Muli, and Ta'u) and those elongate on the ⁸⁷Sr/⁸⁶Sr axis. Mixing components are identified as DMM, HIMU, EM2 and the high ³He/⁴He reservoir, PHEM/FOZO. See Figure 6 for references.

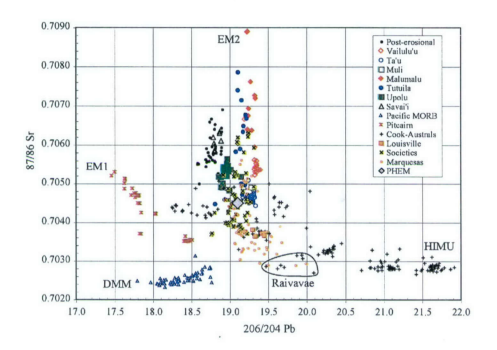


Figure 2-14. Plot showing Sr and Pb isotopic compositions for ocean islands of the Pacific Ocean. Data has been compiled from this study and the GEOROC database. EM2 dominates the spread in composition for the volcanoes Malumalu and Tutuila. Upolu volcano has a significant DMM component and Vailulu'u and Ta'u have been contaminated by HIMU from the Cook-Austral under-plated Pacific lithosphere.

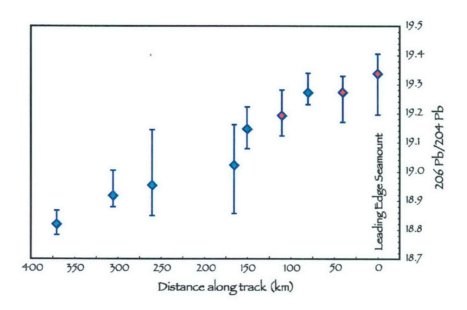


Figure 2-15. Plot showing a systematic increase in ²⁰⁶Pb/²⁰⁴Pb with eastward younging of volcanoes. Distance is measured from the zero-aged leading edge seamount, Vailulu'u. The "oldest" volcano (at a distance of 370 km from Vailulu'u) is Savai'i, though no lavas have been shown to be as old as the theoretical 5 Myr age of the island as suggested from age progression models. High ²⁰⁶Pb/²⁰⁴Pb values are found in EM2 and HIMU; low ²⁰⁶Pb/²⁰⁴Pb values are found in DMM. The increase in ²⁰⁶Pb/²⁰⁴Pb with time is therefore a waning of the DMM component in Samoan lavas.

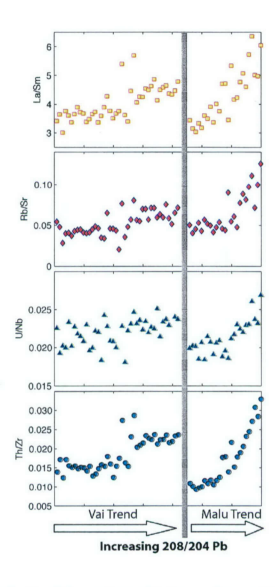


Figure 2-16. Trace element ratios of Samoan lavas, with the more incompatible element in the numerator, showing correlation with ²⁰⁸Pb/²⁰⁴Pb isotopic compositions. The Vai and Malu Trends have been separated into two groups, each sorted by increasing ²⁰⁸Pb/²⁰⁴Pb, and plotted with trace element ratios.

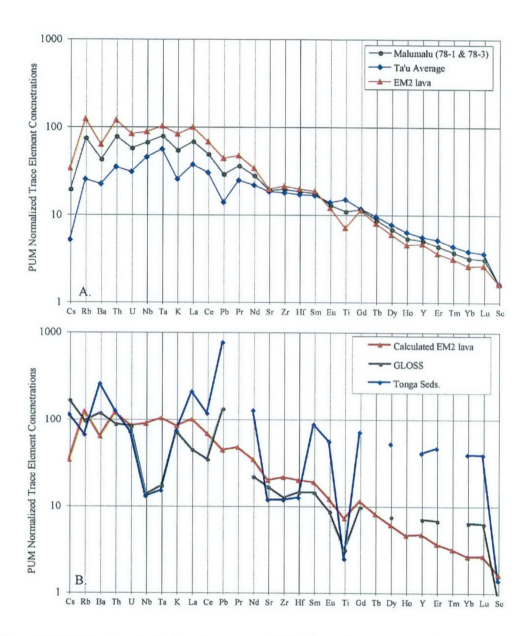


Figure 2-17. Spidergrams in a) show the average of Ta'u lavas, the average of the two most enriched Malumalu lavas, and a calculated EM2 lavas based on extrapolation between Ta'u and Malumalu trace element patterns shown here. All lavas have been corrected for olivine fractionation. In b), the calculated EM2 lava is compared to trace element patterns for globally subducting sediment (GLOSS) and a local Tongan sediment (both from Plank and Langmiur, 1998). Clearly, the trace element patterns between the EM2 lava and sediment are a near-zero match.

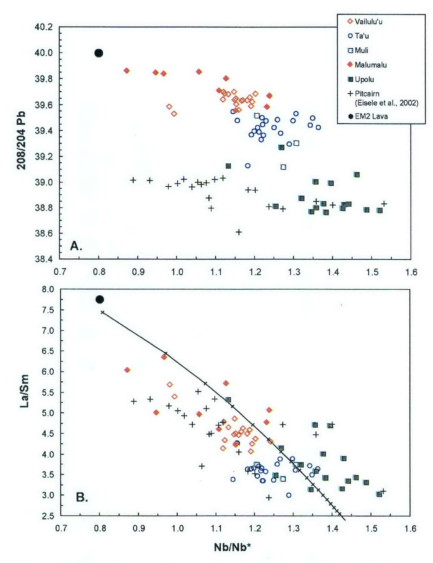


Figure 2-18. Nb/Nb* (calculated as Nb_N/\(Th_N×La_N), as in Eisele et al. [2002]) plotted with a) 208 Pb/ 204 Pb and b) La/Sm, of Samoan lavas and Pitcairn lavas (from Eisele et al., 2002). Pitcairn lavas have little source variation, as seen by a narrow range in 208 Pb/ 204 Pb, but they have a range in Nb/Nb* and La/Sm that is nearly identical to Samoa. This indicates that varying degrees of melting of the same source can provide a wide range of trace element ratios otherwise interpreted to be source variations. The negative correlation in Samoa shows that at small degrees of melting (i.e. high La/Sm and low Nb/Nb*), the enriched component may be preferentially sampled from the mantle. The melting curve is for bacth melting of a mantle with the following concentrations in ppm: Th = 0.032, Nb = 0.457, La = 0.32, Sm = 0.326. D values for these elements are respectively 0.00038, 0.0043, 0.0045, 0.04. Tick marks are every 0.1% melting, increasing toward low La/Sm.

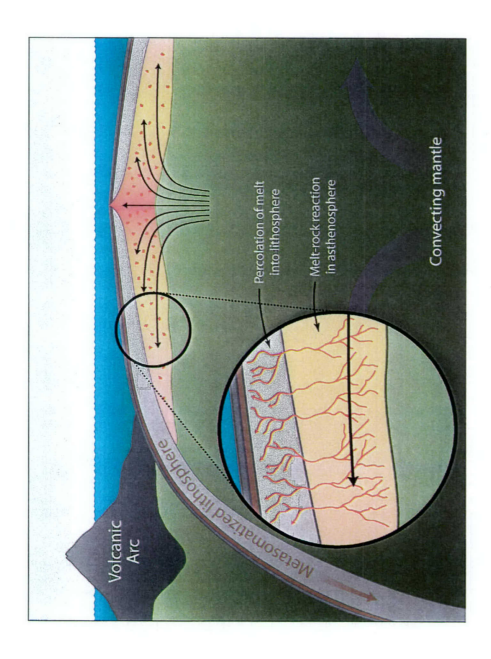


Figure 2-19. Schematic diagram illustrating a working hypothesis for the origin of the EM2 mantle reservoir. Starting 2.5 Ga, small degree (0.5%) batch melts of the primitive upper mantle migrate through the asthenosphere an impregnate the lithosphere. A mixture of depleted lithosphere with 1.1% of the 0.5% batch melts has the trace element pattern required to evolve to the present day Sr, Nd, and Hf isotopic compositions of EM2.

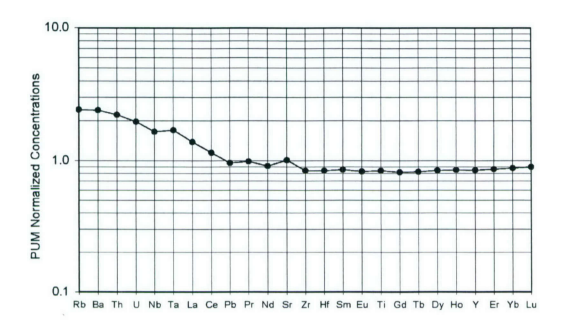


Figure 2-20. Calculated trace element pattern for the EM2 source. At a theoretical 2.5 Ga, a 0.5% batch melt from a primitive upper mantle source has been calculated with a combination of garnet peridotite D values (weighted 72%) and spinel peridotite D values (weighted 28%) from a compilation by Kelemen et al. (2003). Exceptions to Kelemen's D values are as follows: $D^{Rb} = 0.0001$, and $D^{U} = 0.0016$ for both garnet and spinel field melting; for garnet melting, $D^{Th} = 0.00038$, $D^{Zr} = 0.05$, and $D^{Hf} = 0.08$; for spinel melting, $D^{Th} = 0.0011$. Mixing of 1.1% of this melt into a semi-depleted lithosphere results in the trace element pattern shown.

Tables

Table 2-1. 40/39 Argon Ages from Upolu and Savai'i, Western Samoa

Sample Number	Location	Steps Used/ Total Steps	³⁹ Ar Fraction Used	40/39 Total Fusion (my)	Weighted Plateau (my)
1110	Upolu, A'ana	5/7	2.1.00.00/	0.070.000	
U10	Shield Upolu, A'ana	5/7	3.1-98.8%	0.972 ± 0.020	0.933 ± 0.011
U12	Shield	5/6	0-92.8%	2.68 ± 0.03	2.65 ± 0.02
	Savai'i, Manase				
S11	Shield Savai'i, Manase	3/6	4.4-64.7%	0.900±0.122	0.236±0.052
S23	Shield Savai'i, Vanu	5/6	0-91.3%	0.590±0.024	0.386±0.01
91SVK-7	River Shield	-	-	2.05±0.?	-

[•] Step-release heating from 600°-1400°C

 ²⁻sigma errors include measurement uncertainties, and uncertainty in J-value (flux gradient from FCT-3 biotite monitor), but not uncertainty in monitor age.

^{• 91}SVK-7 is a trachyte cobble from the lower Vanu River, analyzed by K. A. Farley

Table 2-2. Sample information and chemical data for Samoan basalts

Volcano Sample	Vailulu'u 63-3	Vailulu'u 63-5	Vailulu'u 63-13	Vailulu'u 64-1	Vailulu'u 68-3	Vailulu'u 68-10	Vailulu'u 68-11
Latitude (°S)	14.218	14.218	14.218	14.260	14.217	14.217	14.217
Longitude (°W)	169.059	169.059	169.059		169.064	169.064	169.064
Water Depth (m)	920	920	920	169.056	780	780	780
Phenocrysts				2630			
rhenocrysts	10% Ol	5% Ol Tr Cpx	Aphyric	3% Ol 2% Cpx	10% OI	Aphyric	60% OI
Major elements	(wt%)	ПСРА		270 Cpx			
SiO ₂	48.29	48.07	47.16	47.71	47.84	48.18	45.61
Al_2O_3	11.88	13.42	14.76	13.38	11.86	14.90	5.64
TiO ₂	2.52	2.67	3.01	2.80	2.55	3.00	1.24
FeO*	10.03	10.19	11.59	10.60	10.51	11.34	11.21
MnO	0.17	0.17	0.18	0.17	0.17	0.19	0.17
CaO	12.78	13.70	12.37	13.48	12.75	12.09	6.23
MgO	10.94	8.26	6.57	8.15	10.68	6.06	28.34
K ₂ O	0.96	0.96	1.28	1.07	0.99	1.13	0.41
Na ₂ O	2.13	2.24	2.72	2.31	2.36	2.76	1.00
P_2O_5	0.29	0.32	0.37	0.33	0.29	0.36	0.14
Mg#	69.58	62.96	54.31	61.72	68.07	52.85	84.13
Trace Elements	(ppm)	110	50	104	205	47	1067
Ni	285	110	59	104	285	47	1067
Cr	879	498	110	455	822	66	2771
V	299	335	339	329	300	351	154
Ga	16	20	20	19	20	18	6
Cu	86	59	58	85	82	68	103
Zn	84	85	95	82	89	94	86
Cs	0.29	0.13	0.28	0.33	0.30	0.38	0.09
Rb	24.7	22.2	33.6	29.6	26.6	24.7	9.3
Ba	216	210	279	236	218	239	90
Th	3.92	3.72	5.15	4.06	4.04	4.67	1.75
U	0.85	0.82	1.07	0.87	0.87	0.98	0.54
Nb	35.41	34.74	45.14	38.97	36.44	42.41	15.57
Ta	2.41	2.32	3.08	2.59	2.50	2.90	1.01
La	28.82	28.95	37.44	31.12	29.29	33.97	13.04
Ce	58.36	58.29	74.35	63.12	58.67	67.35	26.90
Pb	2.41	2.08	2.76	3.04	2.50	2.96	0.66
Pr	6.84	7.00	8.74	7.58	6.86	8.07	3.19
Nd	27.84	29.05	35.38	31.29	29	32.91	12.96
Sr	378	404	470	424	387	434	181
Zr	168	178	218	189	168	202	80
Hf	4.46	4.69	5.76	5.13	4.43	5.45	2.03
Sm	6.45	6.80	7.83	7.33	6.46	7.62	3.01
Eu	2.03	2.13	2.45	2.26	1.99	2.39	0.93
Gd	5.75	6.16	7.09	6.72	5.81	6.83	2.58
Tb	0.88	0.92	1.08	1.02	0.86	1.06	0.40
Dy	4.90	5.17	5.88	5.68	4.83	5.95	2.24
Но	0.87	0.94	1.10	1.01	0.88	1.11	0.41
Y	22.08	24.09	27.50	26.43	22.46	27.93	10.47
Er	2.14	2.32	2.70	2.53	2.13	2.77	0.99
Tm	0.28	0.30	0.35	0.33	0.28	0.36	0.13
Yb	1.60	1.75	2.02	1.91	1.63	2.12	0.74
Lu	0.24	0.25	0.30	0.27	0.24	0.31	0.11
Sc	40.4	42.6	33.4	42.2	40.9	34.0	22.8

^{**} Mg# = molar ratio of MgO/(MgO + 0.85*FeO)

Table 2-2, page 2.

Volcano	Vailulu'u						
Sample	68-28	68-30	70-1	70-2	70-9	71-2	71-11
Latitude (°S)	14.217	14.217	14.208	14.208	14.208	14.337	14.337
Longitude (°W)	169.064	169.064	169.032	169.032	169.032	169.980	169.980
Water Depth (m)	780	780	1130	1130	1130	4170	4170
Phenocrysts	Tr Ol	Aphyric	Aphyric	Tr Cpx	Tr Ol	40% OI	2% OI
					Tr cpx	Tr Cpx	1% Cpx
Major elements	(wt%)				•		
SiO ₂	47.42	48.41	47.57	47.02	47.92	43.85	47.57
Al_2O_3	15.37	12.96	13.63	14.12	14.84	6.37	13.71
TiO ₂	3.62	2.66	2.91	3.01	2.92	1.44	2.75
FeO*	11.49	9.97	11.44	11.88	11.65	10.80	10.64
MnO	0.18	0.17	0.18	0.18	0.18	0.17	0.18
CaO	12.30	14.11	13.05	12.80	11.93	8.07	12.72
MgO	5.01	8.21	7.56	7.02	6.47	27.44	8.60
K ₂ O	1.25	0.96	0.98	1.11	1.05	0.59	1.15
Na ₂ O	2.86	2.27	2.35	2.51	2.70	1.11	2.35
P_2O_5		0.30	0.34	0.35	0.35	0.17	0.33
Mg#	47.77	63.33	58.09	55.34	53.80	84.20	62.90
Trace Elements	(ppm)		0.00		22.00	01.20	02.50
Ni	21	124	57	44	49	1080	135
Cr	50	542	98	72	60	2141	316
V	394	314	338	357	345	177	324
Ga	24	21	21	20	21	9	19
Cu	26	79	82	58	62	138	91
Zn	99	82	92	92	94	77	92
Cs	0.38	0.19	0.31	0.32	0.31	0.19	0.34
Rb	33.1	22.1	25.0	25.8	25.2	16.9	30.4
Ba	292	209	203	245	233	128	247
Th	4.96	3.56	4.20	4.53	4.29	2.31	4.35
U	1.03	0.79	0.93	0.94	0.95	0.49	0.95
Nb	48.08	34.11	37.07	42.81	41.41	20.66	40.53
Та	3.25	2.31	2.50	2.90	2.76	1.38	2.75
La	41.67	27.34	31.03	34.38	32.79	17.10	32.93
Ce	84.11	55.47	64.97	67.72	66.48	34.37	65.58
Pb	2.65	2.36	2.23	2.72	2.64	1.28	2.31
Pr	10.01	6.83	7.57	8.06	7.97	4.04	7.85
Nd	40.82	28.27	31.51	32.96	32.41	16.52	31.64
Sr	521	392	420	453	430	224	434
Zr	222	175	198	199	199	98	186
Hf	5.92	4.72	5.36	5.42	5.24	2.55	5.17
Sm	9.26	6.73	7.50	7.66	7.51	3.68	7.25
Eu	2.81	2.13	2.30	2.38	2.37	1.14	2.25
Gd	8.06	6.06	6.78	6.76	6.77	3.17	6.36
Tb	1.23	0.93	1.02	1.05			0.99
Dy	6.64	5.27	5.67	5.79	5.78	0.48 2.72	5.44
Но	1.20	0.94	1.01	1.06	1.07	0.49	0.98
Y	31.28	24.21	26.38	27.34			
Er	2.96	2.28			27.97	12.57	25.93
Tm	0.38	0.30	2.48	2.56	2.71	1.15	2.39
			0.33	0.34	0.36	0.16	0.32
Yb	2.24	1.68	1.87	1.95	2.07	0.87	1.83
Lu Sc	0.32 32.8	0.25 45.1	0.27 41.9	0.29 38.6	0.31 35.1	0.13 29.8	0.27 39.2

Table 2-2, page 3.

Volcano Sample	Vailulu'u 71-22	Vailulu'u 72-2	Vailulu'u 73-1	Vailulu'u 73-2	Vailulu'u 73-3	Vailulu'u 73-12	Ta'u T10
Latitude (°S)	14.337	14.184	14.214	14.214	14.214	14.214	14.218
Longitude (°W)	169.980	169.894	169.059	169.059	169.059	169.059	169.509
Water Depth(m)	4170	3835	960	960	960	960	-
Phenocrysts	40% OI	Aphyric	Tr Cpx	15% O1	1% O1	15% OI	Tr Ol
			т. ор		Tr Cpx		
Major elements	(wt%)						
SiO ₂	45.21	45.53	47.14	47.42	46.90	47.10	46.60
Al_2O_3	8.13	14.10	13.97	11.29	14.12	12.07	12.41
TiO ₂	1.68	3.31	2.80	2.34	2.87	2.46	3.54
FeO*	10.77	12.13	11.26	10.66	11.49	10.25	12.35
MnO	0.17	0.19	0.18	0.17	0.18	0.18	0.19
CaO	9.99	11.88	12.93	13.18	12.58	12.80	10.98
MgO	21.79	8.56	7.52	11.43	7.71	11.70	10.13
K ₂ O	0.67	1.40	1.37	0.95	1.23	1.15	0.92
Na ₂ O	1.39	2.54	2.50	2.30	2.56	2.02	2.43
P_2O_5	0.20	0.37	0.34	0.26	0.35	0.28	0.45
Mg#	80.93	59.68	58.34	69.23	58.46	70.53	63.24
Trace Elements	(ppm)						
Ni	778	110	79	303	101	220	214
Cr	1687	227	202	840	241	768	579
V	212	367	325	292	328	295	336
Ga	14	23	14	17	22	17	19
Cu	156	79	70	109	66	59	72
Zn	78	102	86	84	89	77	111
Cs	0.21	0.39	0.38	0.11	0.32	0.31	0.12
Rb	19.0	29.3	38.1	24.5	33.2	30.4	19.9
Ba	149	287	347	203	283	282	178
Th	2.62	4.95	6.08	3.90	4.83	4.85	2.77
U	0.56	1.11	1.15	0.83	1.00	0.87	0.73
Nb	24.69	47.47	46.68	34.41	44.50	38.17	35.18
Та	1.64	3.18	3.09	2.31	2.93	2.53	2.52
La	20.16	38.26	44.17	27.56	36.83	36.11	29.20
Ce	40.44	76.30	83.63	54.86	72.25	68.75	62.19
Pb	1.59	2.53	4.79	1.78	2.62	6.23	2.02
Pr	4.73	9.08	9.36	6.38	8.48	7.78	8.14
Nd	19.03	37.20	35.97	26.50	33.93	30.18	34.70
Sr	267	492	473	360	467	398	437
Zr	117	221	212	159	204	177	223
Hf	3.11	5.89	5.77	4.21	5.33	4.89	5.97
Sm	4.36	8.35	7.77	6.15	7.58	6.70	8.35
Eu	1.38	2.60	2.36	1.92	2.36	2.07	2.55
Gd	3.92	7.32	6.77	5.51	6.67	5.95	7.92
Tb	0.60		1.05	0.84	1.03	0.90	1.14
	3.34	6.03	5.64	4.59	5.57	4.96	6.14
Dy	0.59	1.10	1.04	0.86	1.02	0.90	1.11
Но				21.99		23.49	29.41
Y	15.55	29.25	27.27		27.61		
Er	1.42	2.66	2.50	2.02	2.50	2.19	2.78
Tm	0.19	0.35	0.33	0.28	0.33	0.28	0.37
Yb	1.11	2.02	1.90	1.54	1.88	1.64	2.14
Lu Sc	0.16 36.0	0.29 38.0	0.28 39.1	0.23 43.1	0.28 37.3	0.23 42.7	0.32

Table 2-2, page 4.

Volcano Sample	Ta'u T14	Ta'u T16	Ta'u T19	Ta'u T21	Ta'u T22	Ta'u T23	Ta'u T25
Latitude (°S)	14.2163	14.2164	14.2192	14.2160	14.2160	14.2174	14.2163
Longitude (°W)	169.5079	169.5094	169.5037	169.4762	169.4726	169.4648	169.4652
Water Depth (m)	20/ 01	- 01	20/ 01	500/ PI			-
Phenocrysts	2% OI	8% OI	2% OI	50% Plag	Aphyric	Aphyric	25% OI
	Tr Cpx			Tr Ol, Cpx			
Major elements	(wt%)						
SiO_2	46.25	46.71	47.14	48.61	46.51	47.49	45.23
Al_2O_3	10.84	11.27	12.12	17.66	14.47	13.44	8.37
TiO ₂	3.24	3.38	3.52	3.31	4.40	3.85	2.42
FeO*	12.48	11.63	12.07	11.34	12.15	12.39	12.80
MnO	0.19	0.18	0.18	0.19	0.18	0.18	0.19
CaO	10.33	10.91	11.43	8.65	10.51	11.84	8.41
MgO	13.26	12.42	9.73	3.18	7.20	6.77	19.87
K ₂ O	0.82	0.86	0.86	2.07	1.15	0.92	0.66
Na ₂ O	2.23	2.23	2.55	4.07	2.93	2.71	1.78
P_2O_5	0.37	0.39	0.39	0.93	0.52	0.42	0.28
Mg#	69.02	69.13	62.83	37.03	55.41	53.40	76.50
Trace Elements	(ppm)	03.10	02.00	57.05	55.41	55.40	70.50
Ni	396	360	194	9	83	79	723
Cr	943	729	542	5	185	112	1406
V	333	340	345	182	369	373	254
Ga	20	20	20	27	22	22	
Cu	116	32	110	101	70		13 69
Zn	126	102	127	150		105	
					124	123	111
Cs Rb	0.12 18.0	0.09	0.09	0.58	0.19	0.05	0.16
		19.8	18.8	49.1	26.3	16.4	15.9
Ba	164	177	164	406	248	190	126
Th	3.51	2.89	3.24	8.26	4.58	3.80	2.59
U	0.64	0.73	0.73	1.79	1.00	0.71	0.57
Nb	33.26	36.62	33.16	78.63	47.97	38.73	24.03
Та	2.27	2.60	2.30	5.09	3.29	2.64	1.66
La	26.79	29.68	27.08	66.60	37.70	31.06	19.90
Ce	55.65	63.08	56.61	136.19	79.07	65.26	41.81
Pb	5.95	1.82	1.47	3.89	2.25	1.46	1.40
Pr	6.80	8.24	7.06	16.23	9.62	7.98	5.20
Nd	29.62	35.06	31.01	67.87	41.52	35.00	22.45
Sr	377	442	407	754	575	472	294
Zr	205	225	220	460	287	241	154
Hf	5.34	5.83	5.52	10.79	7.40	6.14	3.92
Sm	7.37	8.15	8.07	15.59	10.04	8.68	5.61
Eu	2.41	2.47	2.53	4.69	3.13	2.75	1.78
Gd	7.20	7.64	7.70	13.74	9.38	8.25	5.22
Tb	1.08	1.08	1.15	1.96	1.37	1.21	0.79
Dy	5.61	5.77	6.41	10.77	7.46	6.82	4.27
Но	1.00	1.03	1.17	1.94	1.34	1.21	0.79
Y	25.74	27.91	28.94	50.43	34.20	30.48	20.21
Er	2.39	2.55	2.72	4.58	3.27	2.94	1.85
Tm	0.31	0.33	0.36	0.60	0.42	0.38	0.24
Yb	1.78	1.97	2.04	3.43	2.38	2.10	1.37
Lu	0.25	0.29	0.30	0.51	0.35	0.30	0.19
Sc	32.4	24.2	35.7	17.0	32.7	34.9	26.8

Table 2-2, page 5.

Volcano Sample	Ta'u T27	Ta'u T30	Ta'u T32	Ta'u T33	Ta'u T38	Ta'u T44	Ta'u T45
Latitude (°S)	14.2161	14.2161	14.2163	14.2163	14.2161	14.2564	14.2556
Longitude (°W)	169.4774	169.4774	169.4390	169.4421	169.4774	169.4312	169.4319
Water Depth (m)	-	-	-	-	-	-	-
Phenocrysts	3% OI	3% OI	Aphyric	10% OI	Aphyric	15% OI	3% OI
i nenoci y sta	Tr cpx	1% Cpx	Aphyric	2% Cpx	Aphyric	10% Cpx	370 01
Major elements	(wt%)	170 Cpx		270 CPA		1070 Срх	
		46.41	10.00	47.22	15 21	16 01	47.02
SiO ₂	48.12		48.02	47.32	45.34	46.84	47.93
Al ₂ O ₃	14.17	11.80	13.50	10.61	13.93	10.63	12.81
TiO ₂	3.86	3.57	3.68	3.09	4.64	2.90	3.80
FeO*	11.42	12.07	12.10	11.61	14.00	11.92	11.30
MnO	0.17	0.18	0.18	0.17	0.20	0.18	0.17
CaO	11.54	10.86	11.71	11.54	11.97	10.11	11.67
MgO	6.36	11.52	6.67	12.40	6.11	14.15	8.36
K_2O	1.06	0.83	0.93	0.75	0.93	0.79	0.87
Na ₂ O	2.85	2.36	2.78	2.19	2.51	2.15	2.67
P_2O_5	0.46	0.41	0.42	0.33	0.39	0.33	0.43
Mg#	53.87	66.68	53.62	69.15	47.79	71.34	60.81
Trace Elements	(ppm)						
Ni	95	298	86	288	63	439	162
Cr	290	682	100	988	59	764	549
V	334	313	356	308	489	283	353
Ga	25	19	22	18	24	17	25
Cu	131	102	133	61	120	92	99
Zn	122	121	121	103	121	115	113
Cs	0.09	0.17	0.10	0.11	0.08	0.09	0.14
Rb	21.4	19.0	19.0	15.7	16.7	17.1	18.4
Ba	208	166	180	140	203	140	170
Th	3.86	3.44	3.66	2.83	3.64	3.05	3.38
U	0.79	0.82	0.88	0.58	0.72	0.63	0.82
Nb	39.56	37.63	37.74	28.68	39.88	28.37	35.66
Ta	2.68	2.55	2.56	1.98	2.76	1.91	2.43
			31.24		30.45	23.94	28.63
La	32.43	30.50		23.65			
Ce	67.26	62.61	64.88	50.60	63.59	50.56	60.31
Pb	1.86	1.43	2.32	1.94	1.45	1.90	1.36
Pr	8.24	7.45	7.94	6.38	7.78	6.32	7.44
Nd	35.65	32.06	34.68	27.80	33.84	27.82	32.80
Sr	492	431	469	366	471	356	458
Zr	251	233	236	190	224	199	239
Hf	6.22	5.66	6.11	4.96	5.98	5.08	6.07
Sm	8.85	7.85	8.69	7.09	8.43	7.07	8.49
Eu	2.82	2.50	2.77	2.25	2.69	2.24	2.74
Gd	8.54	7.39	8.30	6.69	8.06	6.70	8.21
Tb	1.29	1.11	1.24	1.02	1.19	1.03	1.23
Dy	7.01	6.03	6.78	5.62	6.48	5.66	6.87
Но	1.28	1.07	1.21	1.01	1.18	1.01	1.24
Y	32.55	27.50	31.25	26.16	29.69	25.28	31.49
Er	3.10	2.62	2.93	2.39	2.78	2.40	2.88
Tm	0.40	0.33	0.38	0.31	0.36	0.31	0.38
Yb	2.28	1.91	2.13	1.74	2.05	1.72	2.16
Lu	0.32	0.28	0.31	0.25	0.29	0.25	0.31
Sc	31.5	32.5	34.6	36.9	40.0	32.4	36.5

Table 2-2, page 6.

Volcano	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u
Sample	T46	T47	T48	T51	T55	74-1	74-4
Latitude (°S)	14.2312	14.2131	14.2176	14.2179	14.2650	14.363	14.363
Longitude (°W)	169.4200	169.4361	169.4202	169.4846	169.4968	169.386	169.386
Water Depth (m)	-	-	-	-	-	2544	2544
Phenocrysts	3% OI	2% O1	1% OI	2% O1	3% O1	25% OI	Aphyric
	2% Cpx	1% Cpx			2% Cpx		
Major elements	(wt%)						
SiO ₂	46.56	48.31	46.13	46.95	48.25	44.09	45.52
Al_2O_3	12.19	11.51	11.75	12.26	12.56	6.67	15.33
TiO ₂	3.51	3.33	3.33	3.55	3.71	1.89	5.76
FeO*	12.46	11.34	12.52	12.02	11.70	12.79	13.92
MnO	0.19	0.17	0.18	0.18	0.18	0.19	0.19
CaO	10.78	12.68	10.17	11.42	12.15	6.98	9.48
MgO	10.38	9.33	12.19	9.82	7.85	25.45	4.83
K ₂ O	0.85	0.71	0.87	0.87	0.73	0.48	1.17
Na ₂ O	2.66	2.26	2.44	2.53	2.48	1.25	3.31
P_2O_5	0.43	0.36	0.41	0.40	0.38	0.22	0.51
Mg#	63.60	63.31	67.13	63.14	58.46	80.67	42.12
Trace Elements	(ppm)					00101	
Ni	230	171	351	194	112	1019	36
Cr	515	654	687	540	331	1655	32
V	323	346	308	341	355	187	472
Ga	20	21	22	22	22	. 13	24
Cu	115	106	85	110	104	69	145
Zn	130	112	120	126	124	103	142
Cs	0.20	0.16	0.05	0.10	0.10	0.13	0.15
Rb	19.2	14.7	17.8	17.5	12.0	10.5	22.3
Ba	176	160	171	177	147	87	237
Th	3.69	3.00	3.55	3.43	2.65	1.51	3.98
U	0.84	0.62	0.74	0.67	0.60	0.42	1.11
Nb	36.94	30.76	35.58	34.21	29.72	18.94	45.78
Ta	2.54	2.14	2.43	2.34	2.04	1.32	3.38
La	30.96	25.08	29.49	27.81	23.86	15.66	37.13
Ce	63.70	53.29	60.93	58.49	51.28	33.14	77.39
Pb	1.66	1.28	1.58	2.05	1.95	1.06	14.60
Pr	7.76	6.62	7.39	7.28	6.52	4.30	9.44
Nd	33.77	29.37	32.39	31.76	29.68	18.16	40.30
Sr	451	396	434	427	427	217	660
Zr	239	199	237	223	214	113	259
Hf	6.18	5.28	5.94	5.72	5.60	3.01	6.79
Sm	8.50	7.47	8.05	8.02	7.95	4.21	9.55
Eu	2.72	2.40	2.57	2.56	2.64	1.28	3.20
Gd	8.17	7.29	7.53	7.72	7.89	3.92	8.68
Tb	1.24	1.09	1.14	1.16	1.20	0.56	1.29
Dy	6.76	6.09	6.28	6.36	6.64	3.04	7.11
Но	1.21	1.09	1.14	1.17	1.21	0.55	1.27
Y	31.09	27.53	28.85	28.91	30.30	14.91	32.34
Er	2.90	2.61	2.74	2.74	2.88	1.40	3.00
Tm	0.37	0.34	0.35	0.36	0.38	0.18	0.40
Yb	2.13	1.91	2.00	1.97	2.10	1.08	2.27
Lu	0.30	0.28	0.29	0.29	0.30	0.16	0.32
Sc	34.0	41.1	31.5	35.8	38.5	18.3	22.5

Table 2-2, page 7.

Volcano Sample	Ta'u 75-10	Muli 79-4	Muli 79-7	Muli 80-23	Malumalu 76-1	Malumalu 76-8	Malumal
Latitude (°S)	14.181	13.916	13.916	14.129	14.569	14.569	14.569
Longitude (°W)	169.304	169.993	169.993	170.062	169.718	169.718	169.718
Water Depth (m)	2675	3484	3484	2368	2785	2785	2785
Phenocrysts	Aphyric	1% Plag	Tr Cpx	Aphyric	Aphyric	2% OI	10% OI
	. ipii jii e	Tr Ol, Cpx	пори	ripingric	ripiijiie	270 01	3% Cpx
Major elements	(wt%)	,					570 Орк
SiO ₂	47.10	47.23	45.88	48.27	47.62	47.04	45.91
Al_2O_3	16.29	17.06	15.99	17.23	14.30	13.69	9.77
TiO ₂	4.02	3.13	3.58	3.36	3.54	3.88	2.74
FeO*	12.56	11.55	12.78	11.05	11.41	11.93	12.20
MnO	0.20	0.22	0.22	0.20	0.16	0.18	0.17
CaO	9.11	8.39	8.93	8.70	12.34	11.86	10.19
MgO	5.16	3.27	4.38	4.21	5.81	6.27	15.84
K ₂ O	1.46	2.09	2.37	1.94	1.69	1.92	1.00
Na ₂ O	3.47	4.70	4.46	4.17	2.75	2.75	1.88
P_2O_5	0.63	2.37	1.40	0.87	0.38	0.49	0.32
Mg#	46.28	37.25	41.82	44.41	51.64	52.43	73.14
Trace Elements	(ppm)	07120	11102		01101	52.15	75.14
Ni	5	4	0	0	46	65	458
Cr	15	1	6	7	94	137	889
V	282	144	168	205	370	356	276
Ga	26	23	27	28	23	22	19
Cu	21	10	6	13	138	72	56
Zn	140	188	191	163	109	121	107
Cs	0.39	0.54	0.68	0.37	0.64	0.51	0.24
Rb	27.6	38.6	54.8	38.4	36.6	44.3	16.9
Ва	264	662	447	409	351	355	187
Th	4.94	6.48	6.82	7.18	6.43	6.41	3.21
U	1.10	1.21	1.74	1.55	1.18	1.30	0.76
Nb	51.26	92.26	77.19	77.72	49.78	56.67	33.01
Та	3.59	5.95	5.28	5.24	3.64	3.93	2.20
La	43.08	107.50	64.09	58.62	51.07	48.43	30.37
Ce	90.50	238.81	137.70	122.09	97.08	95.90	62.55
Pb	3.26	3.80	5.06	4.52	4.32	4.24	2.50
Pr	11.08	29.95	17.21	14.81	11.13	11.17	7.94
Nd	47.98	132.41	77.37	65.12	45.11	46.27	32.42
Sr	617	1901	1010	967	512	544	356
Zr	327	371	492	419	209	276	180
Hf	8.31	9.45	11.73	10.18	6.01	7.23	4.79
Sm	11.62	28.68	18.84	15.63	10.20	10.52	7.19
Eu	3.75	8.77	5.94	4.94	2.98	3.19	2.05
Gd	10.74	23.57	17.17	14.04	9.17	9.49	6.56
Tb	1.60	3.22	2.46	2.04	1.33	1.38	0.91
Dy	8.83	16.70	12.86	10.92	7.08	7.42	4.81
Но	1.59	2.85	2.19	1.88	1.25		0.85
Y	40.24	73.39	56.38	48.07	32.32	1.28 32.16	23.37
Er	3.81	6.52	4.97	4.39	2.97		
	0.50				0.38	2.97	2.06
Tm		0.81 4.39	0.61 3.34	0.56	2.09	0.38	0.27
Yb	2.80			3.07		2.05	1.54
Lu Sc	0.40 20.5	0.62 10.4	0.47 14.1	0.45 15.7	0.31 32.8	0.29 31.0	0.22 24.0

Table 2-2, page 8.

Volcano Sample	Malumalu 76-13	Malumalu 77-1	Malumalu 77-9	Malumalu	Malumalu 78-3	Malumalu	Upolu
				78-1		78-8	U10
Latitude (°S)	14.569	14.701	14.701	14.623	14.623	14.623	13.9283
Longitude (°W)	169.718	169.767	169.767	169.726	169.726	169.726	171.9813
Water Depth (m)	2785	3605	3605	2264	2264	2264	-
Phenocrysts	Aphyric	3% OI	3% OI	25% OI	30% OI	2% O1	Aphyric
		2% Cpx		5% Cpx	5% Cpx	Tr Cpx	
Major elements	(wt%)						
SiO_2	46.35	46.55	46.83	45.54	45.44	47.33	45.46
Al_2O_3	13.55	13.22	13.80	8.94	9.49	14.19	13.21
TiO_2	3.86	3.55	3.58	1.96	2.06	3.29	3.91
FeO*	12.78	12.17	11.08	11.89	11.91	11.45	13.09
MnO	0.20	0.17	0.17	0.17	0.17	0.17	0.19
CaO	11.80	11.90	11.34	8.23	8.41	11.47	11.27
MgO	6.42	7.62	8.45	19.68	18.81	6.93	8.97
K_2O	1.94	1.70	1.81	1.41	1.47	2.05	1.52
Na ₂ O	2.61	2.70	2.59	1.92	1.96	2.74	2.05
P_2O_5	0.49	0.42	0.36	0.27	0.28	0.40	0.34
Mg#	51.30	56.77	61.53	77.63	76.81	55.93	58.97
Trace Elements	(ppm)						
Ni	71	116	154	616	578	97	142
Cr	130	252	391	1289	1175	106	433
V	384	356	370	194	213	342	346
Ga	26	24	21	14	16	21	21
Cu	95	97	62	49	53	79	51
Zn	125	109	108	102	101	104	106
Cs	0.50	0.51	0.55	0.39	0.36	0.60	0.35
Rb	42.8	46.8	47.5	41.8	40.0	51.4	42.8
Ba	341	341	336	255	261	372	341
Th	5.33	5.14	6.59	6.03	5.24	6.26	2.16
U	1.29	1.22	1.24	1.05	1.08	1.36	0.56
Nb	57.53	55.47	53.44	39.02	41.45	58.97	30.02
Ta	3.84	3.70	3.64	2.63	2.71	3.84	2.22
La	48.57	46.34	46.05	39.44	41.66	51.99	23.29
Ce	96.33	90.88	89.34	73.02	77.38	98.76	48.32
Pb	5.18	4.14	4.18	4.19	3.77	4.58	3.11
Pr	11.80	11.05	10.13	7.93	8.85	11.52	5.93
Nd	47.24	43.33	41.30	31.00	32.86	44.15	26.24
Sr	554	530	477	333	361	529	475
Zr	281	250	232	183	193	256	155
Hf	7.11	6.38	6.23	4.71	4.80	6.42	4.41
Sm	10.18	9.15	9.27	6.53	6.56	9.09	6.77
Eu	2.87	2.59	2.72	1.89	1.75	2.51	2.25
Gd	9.10	8.09	8.16	5.81	5.67	7.96	6.60
Tb	1.26	1.12	1.22	0.82	0.77	1.11	1.03
Dy	6.57	5.96	6.52	4.40	4.00	5.84	5.59
Но	1.15	1.04	1.15	0.77	0.70	1.04	1.00
Y	33.12	29.97	29.31	19.57	20.49	30.47	25.19
Er	2.82	2.55	2.72	1.76	1.73	2.53	2.38
Tm	0.37	0.34	0.34	0.23	0.23	0.34	0.30
Yb	2.13	1.92	1.92	1.24	1.32	1.97	1.70
Lu	0.31	0.29	0.28	0.19	0.19	0.29	0.24
Sc	25.0	27.4	31.2	25.2	21.9	28.5	28.0

Table 2-2, page 9.

Volcano Sample	Upolu U12	Upolu U14	Upolu U16	Upolu U19	Upolu U21	Upolu U22	Upolu U24
Latitude (°S)	13.9456	13.9383	13.9355	13.9392	13.8926	13.8922	13.9314
Longitude (°W)	171.9766	171.9805	171.9991	171.9938	171.9640	171.9630	171.9661
Water Depth (m)	-	-	-	-	171.5040	-	-
water Depth (iii)			3% Plag,				_
Phenocrysts	Tr Ol,Plag	2% Plag	Ol	8% Plag	2% Cpx	4% Cpx	5% Plag
i memoery sts	Tr Cpx	2701146	Tr Cpx	2% Cpx	Tr Ol, Plag	2% Ol	Tr Ol, Cp
Major elements	(wt%)		пера	270 СРА	11 01, 1 145	27001	11 01, 00
SiO ₂	48.08	47.51	47.65	47.36	48.24	47.24	49.01
Al_2O_3	14.32	14.53	14.49	15.63	13.88	14.32	15.85
TiO ₂	4.04	4.34	4.03	3.83	3.79	4.00	3.92
FeO*	12.91	13.41	12.20	11.94	12.04	12.84	11.71
MnO	0.19	0.20	0.18	0.17	0.17	0.18	0.18
CaO	10.14	9.94	10.47	11.17	10.64	9.95	9.56
MgO	5.95	5.13	6.30	5.72	7.15	6.98	4.61
K ₂ O	0.96	1.22	1.16	0.99	1.00	1.19	1.35
Na ₂ O	2.90	3.12	2.99	2.74	2.59	2.77	3.19
P_2O_5	0.51	0.60	0.54	0.46	0.49	0.54	0.63
Mg#	49.15	44.51	51.99	50.12	55.46	53.27	
Trace Elements	(ppm)	44.31	31.99	30.12	33.40	33.21	45.22
Ni Ni	72	43	95	73	122	131	43
Cr	137	59	191	162	246		
V	360					211	64
Ga	24	361 24	353	338	354	368	341
Cu	31	38	24 75	22	23 48	22	22
Zn	122	130	112	61 105		92 125	35
Cs	0.09				116		117
Rb	18.0	0.13 26.8	0.15 26.7	0.10	0.40	0.23	0.15
				21.9	24.8	27.7	32.1
Ba	223	255	247	218	220	260	290
Th U	2.64	3.40	2.93	2.59	2.74	3.14	3.79
	0.70	0.90	0.79	0.69	0.77	0.79	0.99
Nb	34.81	45.28	42.59	36.23	35.77	42.88	47.60
Та	2.47	3.21	3.02	2.59	2.54	3.02	3.26
La	30.14	37.48	33.30	29.03	35.27	37.73	40.81
Ce	63.91	80.50	71.31	61.80	70.00	73.99	83.91
Pb	3.32	3.70	3.54	6.98	3.14	3.79	3.93
Pr	8.24	10.11	8.98	7.73	9.26	9.78	10.37
Nd	37.45	44.40	40.23	34.51	41.25	43.19	45.31
Sr	447	535	505	530	479	488	597
Zr	263	313	302	247	253	271	303
Hf	6.85	7.91	7.42	6.35	6.49	6.92	7.51
Sm	9.59	10.93	10.03	8.65	10.12	10.50	10.90
Eu	3.21	3.60	3.28	2.86	3.50	3.55	3.60
Gd	9.35	10.47	9.45	8.27	9.96	10.12	10.35
Tb	1.46	1.59	1.42	1.25	1.48	1.51	1.56
Dy	8.06	8.51	7.76	6.87	7.99	8.34	8.44
Но	1.47	1.56	1.38	1.25	1.45	1.50	1.53
Y	37.16	39.30	35.14	31.35	37.88	38.35	39.08
Er	3.53	3.68	3.29	2.99	3.43	3.54	3.70
Tm	0.46	0.49	0.43	0.38	0.44	0.47	0.47
Yb	2.62	2.68	2.36	2.12	2.46	2.59	2.65
Lu	0.37	0.39	0.34	0.31	0.35	0.38	0.38
Sc	31.1	26.8	29.4	29.4	30.1	28.5	23.9

Table 2-2, page 10.

Sample	Savai'i S11	Savai'i S12	Savai'i S15	Savai'i S16	Savai'i S18	Savai'i S23	Savai'i S25
Latitude (°S)	13.4840	13.4614	13.4561	13.4767	13.4751	13.4469	13.4469
Longitude (°W)	172.3787	172.3654	172.3939	172.3950	172.3929	172.3900	172.3864
Water Depth (m)	172.3767	172.3034	172.3939	172.3930	172.3929	172.3900	
Phenocrysts	3% OI	2% OI	2% OI	Aphyric	Ambumia	2% OI	10/ 01
i lichocrysts	2% Cpx	270 01	Tr Cpx	Aphyric	Aphyric	2% OI	1% O1
14 : 1			перх				
Major elements	(wt%)	46.20		10.17			
SiO ₂	45.95	46.30	45.45	47.16	47.95	45.46	45.57
Al_2O_3	13.27	12.84	12.80	16.43	14.92	11.80	12.00
TiO ₂	3.31	2.96	3.49	3.35	2.70	3.09	3.16
FeO*	12.08	11.91	13.06	11.78	10.66	11.60	11.79
MnO	0.17	0.17	0.18	0.17	0.17	0.17	0.17
CaO	10.54	9.91	7.81	8.45	10.87	9.81	10.07
MgO	9.86	11.86	13.94	6.62	8.23	13.54	13.09
K_2O	1.39	1.35	1.42	1.75	1.28	1.55	0.49
Na ₂ O	2.91	2.24	1.41	3.78	2.87	2.54	3.21
P_2O_5	0.53	0.46	0.44	0.52	0.36	0.44	0.46
Mg#	63.12	67.62	69.12	54.10	61.82	71.00	69.96
Trace Elements	(ppm)						
Ni	193	286	451	86	137	406	383
Cr	415	502	612	77	298	555	568
V	294	271	243	258	271	250	258
Ga	20	18	23	25	22	19	17
Cu	39	44	55	22	57	51	50
Zn	109	106	112	111	85	99	101
Cs	0.31	0.34	0.33	0.54	0.31	0.49	0.52
Rb	34.7	30.4	34.8	53.1	34.4	45.8	116.3
Ba	379	363	489	404	308	430	435
Th	3.87	3.53	3.41	4.43	2.67	3.35	3.46
U	0.78	0.71	0.51	0.91	0.57	0.74	0.72
Nb	44.21	40.01	40.20	46.64	34.71	41.43	42.42
Ta	2.99	2.87	2.89	3.28	2.46	2.85	2.97
La	40.18	34.41	33.75	42.60	29.28	33.20	34.48
Ce	78.56	67.10	66.36	78.71	52.74	66.01	68.38
Pb	4.30	4.06	4.05	5.11	3.44	4.29	3.92
Pr	9.02	7.76	7.76	9.03	6.43	7.75	8.08
Nd	37.22	32.31	33.13	37.01	27.41	32.74	34.15
Sr	581	532	420	680	429	544	564
Zr	213	194	185	251	167	186	189
Hf	5.34	4.97	4.90	6.14	4.27	4.70	4.91
Sm	8.58	7.46	7.84	8.49	6.46	7.72	8.02
Eu	2.81	2.43	2.54	2.84	2.21	2.47	2.60
Gd	8.14	7.09	7.32	8.22	6.74	7.14	7.42
Tb	1.19	1.05	1.06	1.23	1.04	1.05	
Dy	6.37	5.74	5.56	6.76	6.03		1.08
Но	1.11	1.01	0.96			5.45	5.73
Y		25.35		1.24	1.17	0.94	0.98
	28.79		24.13	35.14	36.39	24.49	24.90
Er	2.59	2.39	2.22	2.96	2.97	2.20	2.26
Tm	0.33	0.30	0.28	0.39	0.38	0.27	0.29
Yb	1.82	1.68	1.54	2.19	2.09	1.50	1.60
Lu	0.26	0.24	0.22	0.32	0.33	0.21	0.22

Table 2-3. Major element electron probe data on submarine glasses from Samoa. Normalized to 100%

Sample #	Volcano	SiO2	TiO ₂	Al ₂ O ₃	FeO*	MnO	MgO	CaO	Na ₂ O	K2O	P ₂ O ₅	n
63-13	Vailulu'u	47.223	3.428	14.612	12.744	0.187	5.486	11.147	3.045	1.639	0.489	2
68-03	Vailulu'u	47.454	3.079	14.899	11.726	0.152	6.037	12.039	2.820	1.359	0.436	2
68-34	Vailulu'u	46.896	3.167	14.484	12.175	0.171	6.283	12.376	2.697	1.304	0.447	2
68-35	Vailulu'u	47.208	3.230	14.546	12.739	0.177	5.797	11.912	2.683	1.251	0.456	2
68-36	Vailulu'u	46.954	3.237	14.642	12.239	0.144	6.016	12.136	2.873	1.326	0.433	2
68-37	Vailulu'u	47.071	3.285	14.595	12.647	0.177	5.914	11.937	2.662	1.279	0.432	2
68-38	Vailulu'u	48.142	3.128	14.846	11.631	0.162	5.948	11.528	2.924	1.256	0.435	2
68-43	Vailulu'u	47.054	3.297	14.792	12.297	0.173	5.966	12.045	2.572	1.349	0.456	2
68-44	Vailulu'u	47.011	3.217	14.534	12.444	0.171	6.026	12.028	2.776	1.336	0.456	3
70-01	Vailulu'u	48.043	3.001	14.474	11.821	0.194	6.316	11.996	2.674	1.104	0.378	4
70-02	Vailulu'u	47.548	3.229	14.855	12.204	0.170	5.880	11.668	2.705	1.305	0.437	2
60-02	Vailulu'u	48.094	3.206	14.672	12.364	0.191	5.669	11.317	2.827	1.229	0.433	2
71-03	Vailulu'u	46.986	3.187	14.150	10.839	0.129	7.055	12.911	2.834	1.473	0.437	2
71-04	Vailulu'u	47.034	3.137	13.900	11.492	0.148	6.950	12.722	2.731	1.458	0.429	2
71-05	Vailulu'u	47.735	2.955	14.670	11.278	0.176	6.568	12.212	2.609	1.379	0.419	2
71-06	Vailulu'u	47.631	3.161	14.503	11.581	0.176	5.968	12.676	2.392	1.498	0.414	4
71-09	Vailulu'u	46.874	3.119	14.172	11.304	0.166	7.022	12.716	2.721	1.452	0.453	2
71-10	Vailulu'u	47.149	3.119	14.375	11.286	0.156	6.823	12.458	2.737	1.489	0.409	4
71-11	Vailulu'u	47.492	2.984	15.202	11.487	0.151	6.208	11.908	2.785	1.346	0.438	2
71-13	Vailulu'u	47.680	2.956	15.156	11.521	0.159	6.214	11.923	2.646	1.340	0.406	2
71-13D	Vailulu'u	47.799	2.994	15.288	11.351	0.180	6.188	11.792	2.723	1.300	0.385	4
71-14	Vailulu'u	46.956	3.133	14.127	11.395	0.148	6.994	12.668	2.676	1.460	0.444	2
71-16	Vailulu'u	47.679	3.013	15.199	11.551	0.168	6.013	11.736	2.837	1.351	0.451	3
71-22	Vailulu'u	48.190	2.961	15.655	11.406	0.176	5.540	11.138	3.042	1.490	0.403	4
71-23	Vailulu'u	46.975	3.129	14.136	11.314	0.155	6.959	12.697	2.689	1.483	0.462	2
71-24	Vailulu'u	46.963	3.077	14.133	11.199	0.164	6.979	12.717	2.860	1.457	0.451	2
71-25	Vailulu'u	47.021	3.163	14.108	11.105	0.162	6.987	12.746	2.804	1.480	0.425	2
71-26	Vailulu'u	47.099	3.099	14.144	11.369	0.191	- 6.841	12.510	2.822	1.486	0.441	2
71-27	Vailulu'u	47.094	3.100	14.074	11.273	0.168	6.973	12.581	2.807	1.473	0.457	2
71-28	Vailulu'u	46.886	3.090	14.304	11.446	0.215	696.9	12.483	2.773	1.452	0.382	3
72-02	Vailulu'u	45.953	3.662	15.543	12.494	0.179	5.401	11.380	3.268	1.613	0.508	2
72-04	Vailulu'u	46.292	3.432	14.939	11.785	0.172	6.365	12.222	2.915	1.415	0.463	2
n = number of analyses	fanalyeae											

n = number of analyses *=All Fe reported as FeO

Table 2-3, Page 2

8 15.055 11.808 0.173 4 13.296 11.599 0.157 5 15.072 11.824 0.180 0 13.300 11.869 0.157 1 15.048 11.551 0.160 1 15.08 11.877 0.165 2 15.108 11.582 0.194 5 15.137 11.211 0.176 0 15.122 11.447 0.160 1 15.260 11.335 0.199 1 15.260 11.335 0.199 1 12.550 10.614 0.178 8 12.860 10.628 0.113 8 14.805 11.251 0.168 9 14.836 11.645 0.168 9 14.998 13.116 0.236 14.836 11.251 0.216 9 14.097 14.034 0.216 14.991 12.852 0.231	Volcano SiO2	110_2	Al_2O_3	FeO*	MnO	MgO	CaO	Na ₂ O	K20	P205	u
Vailulu'u 46.842 2.944 13.296 11.599 0.157 Vailulu'u 46.849 3.465 15.072 11.824 0.180 Vailulu'u 46.549 3.000 13.300 11.869 0.183 Vailulu'u 47.342 2.933 15.048 11.551 0.160 Vailulu'u 47.365 2.945 15.016 11.470 0.140 Vailulu'u 47.585 2.955 15.137 11.211 0.176 Vailulu'u 47.603 3.020 15.122 11.447 0.160 Vailulu'u 47.268 2.991 15.260 11.335 0.139 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.766 3.035 14.887 11.251 0.169 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Vailulu'u 48.467 2.68			15.055	11.808	0.173	6.359	12.410	2.794	1.449	0.449	2
Vailulu'u 46.324 3.465 15.072 11.824 0.180 Vailulu'u 46.549 3.000 13.300 11.869 0.183 Vailulu'u 47.342 2.933 15.048 11.551 0.160 Vailulu'u 47.495 2.942 15.108 11.582 0.194 Vailulu'u 47.585 2.942 15.108 11.470 0.140 Vailulu'u 47.585 2.955 15.122 11.417 0.140 Vailulu'u 47.603 3.020 15.122 11.447 0.160 Vailulu'u 48.855 2.527 12.550 10.614 0.176 Vailulu'u 48.855 2.527 12.550 10.618 0.197 Vailulu'u 48.855 2.527 12.550 10.628 0.113 Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 48.214 3.65			13.296	11.599	0.157	8.682	12.103	2.641	1.268	0.468	2
Vailulu'u 46.549 3.000 13.300 11.869 0.183 Vailulu'u 47.342 2.933 15.048 11.551 0.160 Vailulu'u 47.495 2.942 15.089 11.877 0.165 Vailulu'u 47.364 2.985 15.016 11.470 0.140 Vailulu'u 47.585 2.955 15.137 11.211 0.176 Vailulu'u 47.603 3.020 15.122 11.447 0.160 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.855 2.527 12.550 10.628 0.113 Vailulu'u 48.855 2.588 12.860 10.628 0.118 Vailulu'u 47.966 3.035 14.887 11.251 0.164 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.506 3.035 14.988 13.116 0.236 Ta'u 48.409 3.	_	3	15.072	11.824	0.180	6.298	12.159	2.850	1.417	0.412	4
Vailulu'u 47.342 2.933 15.048 11.551 0.160 Vailulu'u 47.213 3.140 15.089 11.877 0.165 Vailulu'u 47.364 2.942 15.108 11.582 0.194 Vailulu'u 47.585 2.942 15.106 11.211 0.176 Vailulu'u 47.603 3.020 15.122 11.447 0.160 Vailulu'u 47.685 2.991 15.260 11.335 0.199 Vailulu'u 47.956 3.268 12.860 10.614 0.178 Vailulu'u 47.956 3.268 14.805 11.958 0.197 Vailulu'u 47.956 3.268 14.805 11.958 0.193 Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 48.467 2.689 13.209 10.765 0.194 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.508 3.	_		13.300	11.869	0.183	8.775	11.905	2.664	1.307	0.448	2
Vailulu'u 47.213 3.140 15.089 11.877 0.165 Vailulu'u 47.495 2.942 15.108 11.582 0.194 Vailulu'u 47.584 2.985 15.106 11.470 0.140 Vailulu'u 47.585 2.955 15.137 11.211 0.176 Vailulu'u 47.268 2.991 15.260 11.335 0.199 Vailulu'u 48.765 2.588 12.860 10.614 0.178 Vailulu'u 48.765 2.588 12.860 10.628 0.113 Vailulu'u 48.765 3.268 14.805 11.958 0.159 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 11.251 0.168 Ta'u 48.467 2.689 11.097 14.034 0.216 Ta'u 48.467 3.035 14.997 14.034 0.216 Ta'u 48.508 3.688 14.991		2	15.048	11.551	0.160	6.028	11.977	2.881	1.639	0.442	3
Vailulu'u 47.495 2.942 15.108 11.582 0.194 Vailulu'u 47.364 2.985 15.016 11.470 0.140 Vailulu'u 47.683 2.955 15.137 11.211 0.176 Vailulu'u 47.683 2.991 15.260 11.335 0.199 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.765 2.588 12.860 10.628 0.113 Vailulu'u 47.956 3.268 14.805 11.558 0.159 Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 14.097 14.034 0.216 Ta'u 48.400 3.728 15.000 12.699 0.235 Ta'u 48.502 3.708 14.991 12.865 0.235 Ta'u 48.502 3.737 15.028 12.815 0.217 Ta'u 48.502 3.708		3	15.089	11.877	0.165	5.905	11.709	2.922	1.510	0.470	2
Vailulu'u 47.364 2.985 15.016 11.470 0.140 Vailulu'u 47.585 2.955 15.137 11.211 0.176 Vailulu'u 47.683 2.991 15.260 11.335 0.199 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.765 2.588 12.860 10.628 0.113 Vailulu'u 47.921 3.377 14.887 11.558 0.159 Vailulu'u 47.966 3.035 14.887 11.645 0.159 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 14.097 14.034 0.216 Ta'u 48.400 3.728 14.097 14.034 0.216 Ta'u 48.508 3.688 14.903 12.865 0.235 Ta'u 48.502 3.734 14.991 12.865 0.220 Ta'u 48.502 3.737 15.028 12.813 0.210 Ta'u 48.502 3.734 1	_		15.108	11.582	0.194	5.972	11.872	2.766	1.636	0.431	2
Vailulu'u 47.585 2.955 15.137 11.211 0.176 Vailulu'u 47.663 3.020 15.122 11.447 0.160 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.765 2.588 12.860 10.628 0.113 Vailulu'u 48.765 3.268 14.805 11.958 0.138 Vailulu'u 47.921 3.377 14.887 11.645 0.159 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.214 3.654 14.988 13.116 0.236 Ta'u 48.400 3.728 14.097 14.034 0.216 Ta'u 48.508 3.688 14.903 12.855 0.220 Ta'u 48.409 3.715 14.991 12.855 0.220 Ta'u 48.508 3.688 14.991 12.855 0.220 Ta'u 48.515 3.737 1	7		15.016	11.470	0.140	6.162	12.073	2.975	1.402	0.412	2
Vailulu'u 47.603 3.020 15.122 11.447 0.160 Vailulu'u 47.268 2.991 15.260 11.335 0.199 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.765 3.268 12.860 10.628 0.113 Vailulu'u 47.921 3.377 14.887 11.645 0.159 Vailulu'u 47.966 3.035 14.887 11.645 0.159 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.214 3.654 14.988 13.116 0.236 Ta'u 48.400 3.728 14.097 14.034 0.216 Ta'u 48.400 3.715 14.991 12.865 0.235 Ta'u 48.508 3.756 14.017 14.165 0.217 Ta'u 48.515 3.737 15.028 12.815 0.220 Ta'u 48.315 3.737 15.028 12.819 0.217 Ta'u 48.315 3.737 15.028	_	(4	15.137	11.211	0.176	5.996	11.976	2.860	1.655	0.449	2
Vailulu'u 47.268 2.991 15.260 11.335 0.199 Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.765 3.588 12.860 10.628 0.113 Vailulu'u 47.956 3.268 14.805 11.958 0.138 Vailulu'u 47.921 3.377 14.887 11.251 0.168 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.214 3.654 14.988 13.116 0.236 Ta'u 48.501 3.728 15.000 12.699 0.220 Ta'u 48.508 3.768 14.901 12.855 0.235 Ta'u 48.409 3.775 14.991 12.85 0.220 Ta'u 48.315 3.737 15.028 12.819 0.210 Ta'u 47.877 3.847 16.455	7	(-)	15.122	11.447	0.160	6.111	12.052	2.695	1.374	0.417	2
Vailulu'u 48.855 2.527 12.550 10.614 0.178 Vailulu'u 48.765 2.588 12.860 10.628 0.113 Vailulu'u 47.921 3.377 14.836 11.645 0.159 Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.513 3.399 14.097 14.034 0.216 Ta'u 48.508 3.788 14.903 12.865 0.231 Ta'u 48.409 3.715 14.991 12.852 0.235 Ta'u 48.409 3.715 14.991 12.85 0.220 Ta'u 48.409 3.775 14.097 14.165 0.217 Ta'u 48.315 3.737 15.028 12.819 0.217 Ta'u 47.877 3.847 16.455 12.320 0.185 Ta'u 46.784 4.073 16.195	7		15.260	11.335	0.199	5.963	11.967	2.995	1.614	0.409	2
Vailulu'u 48.765 2.588 12.860 10.628 0.113 Vailulu'u 47.956 3.268 14.805 11.958 0.138 Vailulu'u 47.921 3.377 14.836 11.645 0.159 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.400 3.728 14.097 14.034 0.216 Ta'u 48.508 3.688 14.907 14.034 0.216 Ta'u 48.409 3.715 14.991 12.855 0.231 Ta'u 48.409 3.715 14.991 12.855 0.235 Ta'u 48.409 3.715 14.991 12.855 0.235 Ta'u 48.409 3.775 14.870 12.819 0.217 Ta'u 48.315 3.737 15.028 12.813 0.217 Ta'u 46.874 4.073 16.155 12.320 0.185 Ta'u 46.782 4.210 16.203 <td< td=""><td></td><td></td><td>12.550</td><td>10.614</td><td>0.178</td><td>8.339</td><td>13.467</td><td>2.242</td><td>0.919</td><td>0.309</td><td>3</td></td<>			12.550	10.614	0.178	8.339	13.467	2.242	0.919	0.309	3
Vailulu'u 47.956 3.268 14.805 11.958 0.138 Vailulu'u 47.921 3.377 14.836 11.645 0.159 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.407 3.539 14.097 14.034 0.216 Ta'u 48.400 3.728 14.903 12.699 0.220 Ta'u 48.409 3.715 14.991 12.852 0.231 Ta'u 48.409 3.715 14.991 12.852 0.235 Ta'u 48.502 3.708 14.870 12.819 0.217 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.209			12.860	10.628	0.113	8.062	13.491	2.270	0.913	0.310	3
Vailulu'u 47.921 3.377 14.836 11.645 0.159 Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.214 3.654 14.988 13.116 0.236 Ta'u 48.400 3.728 14.097 14.034 0.216 Ta'u 48.508 3.688 14.903 12.699 0.220 Ta'u 48.409 3.715 14.991 12.855 0.231 Ta'u 48.409 3.715 14.991 12.852 0.235 Ta'u 48.315 3.778 14.870 12.819 0.217 Ta'u 48.315 3.778 14.870 12.819 0.217 Ta'u 47.857 3.762 16.359 12.515 0.223 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.096 16.323 12.9			14.805	11.958	0.138	5.427	11.164	3.220	1.602	0.463	3
Vailulu'u 47.966 3.035 14.887 11.251 0.168 Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.214 3.654 14.988 13.116 0.236 Ta'u 48.508 3.728 15.000 12.699 0.220 Ta'u 48.508 3.688 14.901 12.865 0.231 Ta'u 48.409 3.715 14.991 12.852 0.235 Ta'u 48.502 3.708 14.870 12.819 0.217 Ta'u 48.502 3.737 15.028 12.813 0.210 Ta'u 47.877 3.847 16.455 12.320 0.185 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.196		6	14.836	11.645	0.159	5.708	11.546	2.941	1.450	0.416	3
Vailulu'u 48.467 2.689 13.209 10.765 0.197 Ta'u 48.214 3.654 14,988 13.116 0.236 Ta'u 48.508 3.728 15.000 12.699 0.220 Ta'u 48.508 3.688 14,903 12.865 0.231 Ta'u 48.409 3.715 14,991 12.852 0.235 Ta'u 48.502 3.708 14.870 12.819 0.217 Ta'u 48.502 3.737 15.028 12.813 0.210 Ta'u 47.877 3.847 16.455 12.320 0.185 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.787 4.210 16.203 12.953 0.211 Ta'u 46.782 4.210 16.203 12.953 0.213 Ta'u 46.782 4.210 16.203 12.953 0.196			14.887	11.251	0.168	6.199	12.108	2.706	1.291	0.388	3
Ta'u 48.214 3.654 14,988 13.116 0.236 Ta'u 45.513 5.399 14,097 14.034 0.216 Ta'u 48.508 3.688 14,903 12.865 0.220 Ta'u 48.409 3.715 14,991 12.852 0.231 Ta'u 48.502 3.736 14,017 14.165 0.220 Ta'u 48.502 3.737 15.028 12.819 0.217 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.787 3.842 16.475 11.959 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.211			13.209	10.765	0.197	7.676	13.297	2.388	0.982	0.329	3
Ta'u 45.513 5.399 14.097 14.034 0.216 Ta'u 48.508 3.728 15.000 12.699 0.220 Ta'u 48.409 3.715 14.991 12.852 0.231 Ta'u 48.502 3.736 14.017 14.165 0.220 Ta'u 48.502 3.737 15.028 12.819 0.217 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 47.857 3.762 16.359 12.515 0.223 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.196		_	14.988	13.116	0.236	3.984	8.720	4.147	1.953	0.988	2
Ta'u 48.400 3.728 15.000 12.699 0.220 Ta'u 48.508 3.688 14.903 12.855 0.231 Ta'u 45.473 5.356 14.017 14.165 0.220 Ta'u 48.502 3.708 14.870 12.819 0.217 Ta'u 48.315 3.737 15.028 12.813 0.210 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.196		2	14.097	14.034	0.216	5.537	10.951	2.784	1.007	0.462	2
Ta'u 48.508 3.688 14,903 12.865 0.231 Ta'u 48.409 3.715 14,991 12.852 0.235 Ta'u 48.502 3.736 14,017 14,165 0.220 Ta'u 48.315 3.737 15.028 12.819 0.217 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 47.857 3.762 16.359 12.515 0.223 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.096 16.323 12.928 0.196		3	15.000	12.699	0.220	3.999	8.854	4.157	1.942	1.002	2
Ta'u 48.409 3.715 14.991 12.852 0.235 Ta'u 45.473 5.356 14.017 14.165 0.220 Ta'u 48.502 3.737 15.028 12.819 0.217 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 47.857 3.762 16.359 12.515 0.223 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.096 16.323 12.928 0.196		3	14.903	12.865	0.231	3.987	8.752	4.124	1.946	0.995	2
Ta'u 45.473 5.356 14.017 14.165 0.220 Ta'u 48.502 3.708 14.870 12.819 0.217 Ta'u 47.977 3.847 15.028 12.320 0.185 Ta'u 47.857 3.762 16.359 12.515 0.223 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.724 4.096 16.323 12.928 0.196		3	14.991	12.852	0.235	4.030	8.715	4.151	1.924	0.977	2
Ta'u 48.502 3.708 14.870 12.819 0.217 Ta'u 48.315 3.737 15.028 12.813 0.210 Ta'u 47.877 3.847 16.455 12.320 0.185 Ta'u 46.874 4.073 16.119 12.920 0.203 Ta'u 48.175 3.842 16.475 11.959 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.724 4.096 16.323 12.928 0.196	_	5	14.017	14.165	0.220	5.556	11.096	2.668	0.991	0.459	2
Ta'u 48.315 3.737 15.028 12.813 0.210 Ta'u 47.977 3.847 16.455 12.320 0.185 Ta'u 47.857 3.762 16.359 12.515 0.223 Ta'u 46.874 4.073 16.119 12.920 0.209 Ta'u 48.175 3.842 16.475 11.959 0.211 Ta'u 46.782 4.210 16.203 12.953 0.223 Ta'u 46.724 4.096 16.323 12.928 0.196		3	14.870	12.819	0.217	4.024	8.786	4.197	1.916	0.960	2
Ta'u 47.977 3.847 16.455 12.320 0.185 4 Ta'u 47.857 3.762 16.359 12.515 0.223 4 Ta'u 46.874 4.073 16.119 12.920 0.209 5 Ta'u 48.175 3.842 16.475 11.959 0.211 5 Ta'u 46.782 4.210 16.203 12.953 0.223 5 Ta'u 46.724 4.096 16.323 12.928 0.196 5		3	15.028	12.813	0.210	4.032	8.858	4.123	1.914	0.971	2
Ta'u 47.857 3.762 16.359 12.515 0.223 4 Ta'u 46.874 4.073 16.119 12.920 0.209 5 Ta'u 48.175 3.842 16.475 11.959 0.211 5 Ta'u 46.782 4.210 16.203 12.953 0.223 5 Ta'u 46.724 4.096 16.323 12.928 0.196 5		3	16.455	12.320	0.185	4.920	8.330	3.775	1.479	0.713	2
Ta'u 46.874 4.073 16.119 12.920 0.209 5 Ta'u 48.175 3.842 16.475 11.959 0.211 5 Ta'u 46.782 4.210 16.203 12.953 0.223 5 Ta'u 46.724 4.096 16.323 12.928 0.196 5		3	16.359	12.515	0.223	4.941	8.315	3.843	1.480	0.706	2
Ta'u 48.175 3.842 16.475 11.959 0.211 5 Ta'u 46.782 4.210 16.203 12.953 0.223 5 Ta'u 46.724 4.096 16.323 12.928 0.196 5		4	16.119	12.920	0.209	5.320	8.657	3.813	1.376	0.639	2
5 Ta'u 46.782 4.210 16.203 12.953 0.223 5 7 Ta'u 46.724 4.096 16.323 12.928 0.196 5		3	16.475	11.959	0.211	5.011	8.420	3.703	1.467	0.736	2
7 Ta'u 46.724 4.096 16.323 12.928 0.196 5		4	16.203	12.953	0.223	5.287	8.663	3.666	1.375	0.640	2
		4	16.323	12.928	0.196	5.324	8.734	3.663	1.378	0.635	2
75-8 Ta'u 47.096 4.030 16.311 12.661 0.222 5.258		4	16.311	12.661	0.222	5.258	8.687	3.748	1.361	0.627	2

Sample #	Volcano	SiO2	TiO2	Al ₂ O ₃	FeO*	MnO	MgO	CaO	Na ₂ O	K2O	P2O5	u
75-12	Ta'u	47.820	3.838	16.317	12.280	0.220	4.850	8.447	3.956	1.562	0.710	
75-13	Ta'u	46.771	4.691	14.903	13.400	0.235	4.961	9.076	3.794	1.474	0.695	
75-15	Ta'u	47.671	3.924	16.476	12.460	0.215	5.017	8.547	3.582	1.460	0.648	7
75-16	Ta'u	47.054	4.064	16.334	12.966	0.222	5.248	8.605	3.541	1.381	0.583	
75-17	Ta'u	46.752	4.726	15.082	13.335	0.247	4.925	9.039	3.689	1.489	0.716	, ,
75-18	Ta'u	46.895	4.097	16.234	12.755	0.198	5.329	8.741	3.811	1.352	0.588	
76-1	Malumalu	46.986	3.831	14.854	12.706	0.179	5.169	10.602	3.262	1.945	0.467	14
76-2	Malumalu	47.471	3.689	15.298	12.818	0.187	4.710	9.971	3.444	2.012	0.400	
76-3	Malumalu	46.960	3.952	14.469	12.521	0.199	5.455	10.813	3.303	1.830	0.498	(4
76-4	Malumalu	46.786	3.963	14.467	12.713	0.179	5.502	10.826	3.222	1.865	0.477	7
76-5	Malumalu	48.855	3.407	15.699	12.653	0.210	5.447	8.873	3.092	1.346	0.418	(-)
9-92	Malumalu	47.388	3.576	15.274	12.282	0.166	5.151	10.393	3.160	2.195	0.416	
7-97	Malumalu	47.544	3.718	15.316	12.705	0.220	4.736	10.050	3.362	1.963	0.386	(4
8-92	Malumalu	46.737	3.892	14.358	12.583	0.187	5.585	11.002	3.346	1.824	0.485	. 4
76-10	Malumalu	47.377	3.982	15.507	12.673	0.191	4.458	9.716	3.464	2.098	0.534	
76-11	Malumalu	47.393	3.608	15.316	12.639	0.199	4.920	10.278	3.389	1.893	0.364	(4
76-12	Malumalu	46.948	3.957	14.504	12.546	0.197	5.486	10.818	3.180	1.869	0.495	(4
76-13	Malumalu	46.869	3.958	14.567	12.434	0.188	5.579	10.875	3.240	1.839	0.451	
76-14	Malumalu	47.499	3.611	15.355	12.479	0.189	4.997	10.163	3.203	2.100	0.404	
9-11	Malumalu	47.023	3.764	14.467	12.561	0.209	5.598	10.995	3.009	1.957	0.417	(4
78-2	Malumalu	48.530	3.255	16.109	10.645	0.196	4.957	10.164	3.214	2.450	0.481	(4
78-10	Malumalu	48.627	3.252	15.758	10.580	0.163	4.965	10.107	3.491	2.612	0.444	(4
79-2	Muli	44.232	5.043	15.262	11.764	0.229	5.401	10.162	4.357	2.520	1.030	
79-3	Muli	44.495	5.024	15.139	11.580	0.209	5.271	10.135	4.587	2.550	1.009	(4
79-5	Muli	48.855	3.153	16.510	10.307	0.243	3.822	8.401	5.013	2.580	1.117	(4
8-62	Muli	48.850	3.143	16.528	10.319	0.228	3.955	8.269	4.962	2.555	1.190	
79-18	Muli	48.876	3.046	16.681	10.241	0.255	3.813	8.022	5.319	2.661	1.086	(4
80-1	Muli	46.873	4.430	15.752	12.337	0.250	4.696	8.808	4.215	1.821	0.820	, ,
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Table 2-4. Isotopic compositions of Samoan lavas.

Volcano	Sample	86Sr/87Sr	143Nd/144Nd	206Pb/204Pb	$^{207}\mathrm{Pb}/^{204}\mathrm{Pb}$	208Pb/204Pb	³He∕⁴He	cc/gm		$^{187}Os^{188}Os$	Os (ppb)
Vailulu'u	63-3	0.705453	0.512745	19.350	15.619	39.698				0.1275	0.261
Vailulu'u	63-5	0.705299	0.512746	19.337	15.604	39.604	1				
Vailulu'u	63-13	0.705520	0.512716	19.352	15.624	39.699	,	1		,	
Vailulu'u	64-1	0.705303	0.512764	19.363	15.606	39.623	•	,		,	•
Vailulu'u	68-3	0.705396		19.337	15.621	39.651	10.04(2)	1.16E-07	ol, gl		
Vailulu'u	68-10	0.705373	0.512760	19.325	15.610	39.560)		
Vailulu'u	68-11	0.705594	0.512724	19.357	15.623	39.685	1	i		0.1280	2.039
Vailulu'u	68-28	0.705419	0.512743	19.337	15.598	39.648	1	i			•
Vailulu'u	68-30	0.705215	0.512734	19.332	15.615	39.589	1			,	
Vailulu'u	70-1	0.705371	0.512768	19.371	15.608	39.641	1	ī		,	,
Vailulu'u	70-2	0.705359	0.512738	19.405	15.618	39.633	1	1			•
Vailulu'u	6-07	0.705352	0.512753	19.386	15.619	39.683	8.05	5.78E-08	-	,	1
Vailulu'u	71-2	0.705943	0.512743	19.332	15.605	39.680	9.48 (2)	1.76E-07	ol, gl	0.1300	1.357
Vailulu'u	71-11	0.705394	0.512743	19.353	15.603	39.630	9.85	5.48E-07	lo		1
Vailulu'u	71-22	0.705473	0.512747	19.358	15.602	39.635	9.64	2.99E-08	10		1
Vailulu'u	72-2	0.705395	0.512740	19.365	15.604	39.657					1
Vailulu'u	73-1	0.706720	0.512669	19.215	15.617	39.585	8.1 (3)	2.09E-08	lg.	i	,
Vailulu'u	73-2	0.705424	0.512742	19.329	15.621	39.616	9.28	1.07E-08	10		ı
Vailulu'u	73-3	0.705616	0.512711	19.328	15.602	39.638	•	,			1
Vailulu'u	73-12	0.706653	0.512686	19.195	15.600	39.529	,			0.1288	0.105
Ta'u	T10	0.704657	0.512789	19.291	15.623	39.497	•	•			,
Ta'u	T14	0.704591	0.512806	18.934	15.590	39.131	•				ı
Ta'u	T16	0.704605	0.512818	19.290	15.601	39.426	15.13	5.73E-09	10	0.1294	0.279
Ta'u	T19	0.704582	0.512790	19.299	15.600	39.448	,	,			1
Ta'u	T21	0.704751	0.512796	19.313	15.594	39.479	,				1
Ta'u	T22	0.704701	0.512773	19.314	15.606	39.485		×		1	1
Ta'u	T23	0.704706	0.512796	19.288	15.614	39.476	,			1	1
Ta'u	T25	0.704708	0.512790	,	,		13.26	1.05E-07	lo	0.1290	0.206
Ta'u	T27	0.704561	0.512806	19.304	15.604	39.500					,
Ta'u	T30	0.704528	0.512822	19.305	15.594	39.382		·		0.1364	0.080
Ta'u	T32	0.704588	0.512797	19.208	15.581	39.332	•				,
Ta'u	T33	0.704736	0.512780	19.284	15.596	39.445	16.62	4.35E-09	lo		
Ta'u	T38	0.704651	0.512785	19.328	15.617	39.532	,			,	
Ta'u	T44	0.705086	0.512755	19.246	15.606	39.549	14.86	9.61E-09	lo	î	1
Ta'u	T45	0.704434	0.512816	19.337	15.595	39.425	í			1	•
Ta'u	T46	0.704676	0.512795	19.231	15.584	39.369	i				1
Ta'u	T47	0.704504	0.512800	19.253	15.595	39.366	,			,	,

Table 2-4. Page 2

Volcano	Sample	86Sr/87Sr	PN41/PN641	206Pb/204Pb	207 Pb/204 Pb	$^{208}Pb/^{204}Pb$	³He/⁴He	cc/gm		sO ₈₈₁ /sO ₂₈₁	Os (ppb)
Ta'u	T48	0.704664	0.512789	19.249	15.595	39.397				0.1351	0.062
Ta'u	T51	0.704614	0.512794	19.276	15.591	39.425		•			
Ta'u	T55	0.704545	0.512815	19.178	15.594	39.298		•			ï
Ta'u	74-1	0.704686	0.512786	19.29	15.599	39.443	17.97	4.97E-08	10	0.1291	1.397
Ta'u	74-4	0.704815	0.512784	19.314	15.601	39.477				,	1
Ta'u	75-10	0.704533	0.512792	19.266	15.597	39.39	•	,		¢	•
Muli	79-4	0.704904	0.512730	19.279	15.617	39.517	,			•	1
Muli	7-67	0.704524	0.512812	19.122	15.581	39.122	,				
Muli	80-23	0.704914	0.512767	19.177	15.591	39.305	i	•		•	
Malumalu	76-1	0.707192	0.512637	19.338	15.636	39.847	,			•	
Malumalu	8-92	0.706374	0.512667	19.294	15.633	39.710		•		•	
Malumalu	6-92	0.706745	0.512669	19.245	15.596	39.555	15.89	2.12E-08	lo	•	•
Malumalu	76-13	0.706395	0.512680	19.237	15.600	39.584		•		•	·
Malumalu	77-1	0.706930	0.512663	19.251	15.619	39.669	13.45	3.30E-08	10	•	
Malumalu	77-9	0.707260	0.512579	19.331	15.635	39.853	10.56	1.92E-08	10		
Malumalu	78-1	0.708901	0.512521	19.237	15.647	39.862	8.09	9.61E-08	lo	0.1293	0.130
Malumalu	78-3	0.708886	0.512511	19.230	15.641	39.840	8.22	9.15E-08	10	0.1288	0.427
Malumalu	78-8	0.707614	0.512580	19.276	15.633	39.803					1
Upolu	010	0.705365	0.512774	19.044	15.582	39.067	1	,		0.1407	0.013
Upolu	U12			18.889	15.554	38.772	•				1.
Upolu	U14			18.878	15.560	38.767	,			•	,
Upolu	010	0.705171	0.512883	18.881	15.559	38.787	,	,			
Upolu	010	0.705278	0.512870	18.917	15.569	38.832	,	,			,
Upolu	U21	0.705011		18.901	15.561	38.814	Y	,			
Upolu	U22			18.912	15.563	38.802	1	•		0.1509	0.022
Upolu	U24	0.705191	0.512854	18.955	15.569	38.875	¥	,		•	
Savaili	S11	0.706195	0.512693	18.782	15.604	38.995	•	,		0.1299	0.107
Savai'i	S12	,		18.799	15.603	39.002	·				
Savai'i	S15	0.706039	0.512686	18.793	15.610	39.022					ř
Savai'i	816	0.706296	0.512705	18.865	15.595	39.089		,		•	•
Savaili	818	0.706110	0.512730	18.884	15.596	39.118					
Savai'i	S23			18.795	15.599	38.985	ì	1.		0.1270	0.491
Savai'i	S25	0.705848	0.512706	18.797	15.600	38.982				0.1353	0.034

Volcanic Series Lat. (°S) Long. (°W) Major elements (w SiO ₂ Al ₂ O ₃ TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _i Ni Cr	48.06 15.20 4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm)	Fagaloa 13.8533 171.6582 45.65 13.98 2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	Fagaloa 13.9422 171.5848 45.78 15.79 3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65 52.31	Fagaloa 13.8453 171.7093 48.28 15.32 4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63 47.85	U 39 F Fagaloa 13.8512 171.7033 47.33 13.86 5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	U 40 F Fagaloa 13.8597 171.6523 47.53 14.00 3.88 12.50 0.17 10.26 7.45 0.97 2.75	U 41 F Fagaloa 13.8648 171.6436 46.01 13.87 2.96 12.67 0.18 10.77 10.00 0.94
Long. (°W) Major elements (w SiO ₂ Al ₂ O ₃ TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p)	171.6886 48.06 15.20 4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	45.65 13.98 2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	13.9422 171.5848 45.78 15.79 3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65	13.8453 171.7093 48.28 15.32 4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63	13.8512 171.7033 47.33 13.86 5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	13.8597 171.6523 47.53 14.00 3.88 12.50 0.17 10.26 7.45 0.97	13.8648 171.6436 46.01 13.87 2.96 12.67 0.18 10.77 10.00 0.94
Long. (°W) Major elements (w SiO ₂ Al ₂ O ₃ TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p)	171.6886 48.06 15.20 4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	45.65 13.98 2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	171.5848 45.78 15.79 3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65	48.28 15.32 4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63	47.33 13.86 5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	171.6523 47.53 14.00 3.88 12.50 0.17 10.26 7.45 0.97	171.6436 46.01 13.87 2.96 12.67 0.18 10.77 10.00 0.94
Major elements (w SiO ₂ Al ₂ O ₃ TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p ₁	48.06 15.20 4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm)	45.65 13.98 2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	45.78 15.79 3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65	48.28 15.32 4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63	47.33 13.86 5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	47.53 14.00 3.88 12.50 0.17 10.26 7.45 0.97	46.01 13.87 2.96 12.67 0.18 10.77 10.00 0.94
SiO ₂ Al ₂ O ₃ TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p ₁	48.06 15.20 4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm)	13.98 2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	15.79 3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65	15.32 4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63	13.86 5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	14.00 3.88 12.50 0.17 10.26 7.45 0.97	13.87 2.96 12.67 0.18 10.77 10.00 0.94
Al ₂ O ₃ TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p ₁	15.20 4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm)	13.98 2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	15.79 3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65	15.32 4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63	13.86 5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	14.00 3.88 12.50 0.17 10.26 7.45 0.97	13.87 2.96 12.67 0.18 10.77 10.00 0.94
TiO ₂ FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _j	4.24 12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	2.92 12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	3.65 12.84 0.19 9.82 6.72 0.56 4.00 0.65	4.28 12.05 0.17 9.37 5.27 1.41 3.22 0.63	5.09 12.25 0.16 10.47 5.80 1.44 2.92 0.68	3.88 12.50 0.17 10.26 7.45 0.97	2.96 12.67 0.18 10.77 10.00 0.94
FeO* MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _j	12.01 0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	12.82 0.18 10.27 10.24 0.91 2.59 0.44 62.63	12.84 0.19 9.82 6.72 0.56 4.00 0.65	12.05 0.17 9.37 5.27 1.41 3.22 0.63	12.25 0.16 10.47 5.80 1.44 2.92 0.68	12.50 0.17 10.26 7.45 0.97	12.67 0.18 10.77 10.00 0.94
MnO CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _j	0.16 9.73 5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	0.18 10.27 10.24 0.91 2.59 0.44 62.63	0.19 9.82 6.72 0.56 4.00 0.65	0.17 9.37 5.27 1.41 3.22 0.63	0.16 10.47 5.80 1.44 2.92 0.68	0.17 10.26 7.45 0.97	0.18 10.77 10.00 0.94
CaO MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _j	9.73 5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	10.27 10.24 0.91 2.59 0.44 62.63	9.82 6.72 0.56 4.00 0.65	9.37 5.27 1.41 3.22 0.63	10.47 5.80 1.44 2.92 0.68	10.26 7.45 0.97	10.77 10.00 0.94
MgO K ₂ O Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _j	5.38 1.37 3.22 0.63 48.44 pm) 72 105 312	10.24 0.91 2.59 0.44 62.63	6.72 0.56 4.00 0.65	5.27 1.41 3.22 0.63	5.80 1.44 2.92 0.68	7.45 0.97	10.00 0.94
$egin{array}{l} K_2^{-}O \\ Na_2O \\ P_2O_5 \\ Mg\# \\ \emph{Trace Elements (p)} \\ Ni \end{array}$	1.37 3.22 0.63 48.44 pm) 72 105 312	0.91 2.59 0.44 62.63	0.56 4.00 0.65	1.41 3.22 0.63	1.44 2.92 0.68	0.97	0.94
Na ₂ O P ₂ O ₅ Mg# Trace Elements (p _j Ni	3.22 0.63 48.44 pm) 72 105 312	2.59 0.44 62.63	4.00 0.65	3.22 0.63	2.92 0.68	0.97	
P ₂ O ₅ Mg# Trace Elements (p _j Ni	0.63 48.44 pm) 72 105 312	0.44 62.63 249	0.65	0.63	0.68		216
Mg# Trace Elements (p _j Ni	48.44 pm) 72 105 312	62.63 249					2.16
Trace Elements (p) Ni	72 105 312	249	32.31	47.00	10 92	0.48	0.45
Ni	72 105 312				49.82	55.54	62.35
	312		72	72	70	1.40	240
	312	389	72 75	73 98	79 73	140	248
V	312	272	293			220	411
	25		23	311	341	316	280
Ga Cu	71	20 92	38	25	25	22	21
	144	123		69	120	101	98
Zn Cs	0.20	0.40	134	150	154	130	124
Rb	32.3	0.40	0.54	0.32	0.47	0.14	0.23
Ba	311	23.8 259	38.2	33.0	38.2	21.2	23.8
Th	3.79	3.17	341	292	305	203	272
Ü	1.07	0.76	5.27 1.12	3.77 1.07	5.08	2.64	3.27
Nb	52.18	40.16	52.88	51.82	1.26 58.00	0.73	0.79
Та	3.26	2.48	3.07	3.19	38.00	35.35	40.33
La	45.10	31.13	49.32	41.43	3.60	2.23	2.50
Ce	96.38	66.54	102.63	93.74	48.93	27.74	32.16
Pb	2.82	2.40	3.72	2.98	105.67 3.48	65.00	67.97
Pr	13.21	8.21	12.34	12.49	13.99	1.70	2.27
Nd	53.08	31.72	46.48	50.08	55.92	9.03 38.14	8.43 32.80
Sr	683	521	699	664	631	494	543
Zr	324	179	243	322	335	264	183
Hf	7.79	4.35	5.71	7.71	8.13	6.46	4.51
Sm	11.24	6.62	9.25	10.61	11.76	8.76	6.82
Eu	3.60	2.11	2.89	3.36	3.66	2.79	2.18
Gd	10.52	6.11	8.19	9.57	10.63	8.18	6.33
Tb	1.46	0.87	1.14	1.34	1.46	1.17	0.89
Dy	8.11 .	4.82	6.25	7.37	7.97	6.56	4.95
Ho	1.49	0.88	1.13	1.33	1.42	1.21	0.90
Y	39.25	21.40	27.44	31.82	34.83	28.89	21.98
Er	3.69	2.17	2.75	3.21	3.44	2.94	2.23
Tm	0.47	0.29	0.36	0.42	0.44	0.39	0.29
Yb	2.69	1.69	2.10	2.42	2.53	2.25	1.71
Lu	0.38	0.24	0.30	0.34	0.35	0.32	0.24
Sc	22.2	22.7	19.0	21.9	25.3	25.9	24.0
⁸⁶ Sr/ ⁸⁷ Sr	0.705361	0.705391	0.705644		0.705180	0.704904	0.705439
143Nd/144Nd	0.512874	0.512783	0.512773			0.512907	0.512777
²⁰⁶ Pb/ ²⁰⁴ Pb	18.944	18.914	18.961	18.940	19.143	18.905	18.918
²⁰⁷ Pb/ ²⁰⁴ Pb	15.580	15.582	15.603	15.576	15.606	15.568	15.584
²⁰⁸ Pb/ ²⁰⁴ Pb	38.835	38.996	39.130	38.827	39.273	38.798	39.009

^{*}See Appendix 1 for analytical techniques. ** Mg# = molar ratio of MgO/(MgO + 0.85*FeO)

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Volcano	Upolu	Savai'i	Savai'i	Savai'i	Savai'i	Savai'i	Savai'i PE
Sample	U 43 F	S 36 F	S 44 F	S 45 F	S 46 F	S 47 F	S 10 M
Volcanic							
Series	Fagaloa	Fagaloa	Fagaloa	Fagaloa	Fagaloa	Fagaloa	Mulifanua
Lat. (°S)	13.8895	13.2642	13.2662	13.2662	13.2642	13.2637	13.3058
Long. (°W)	171.5614	172.3815	172.3677	172.3674	172.3813	172.3841	172.6411
Major elemen	ts (wt%)						
SiO ₂	46.66	45.93	47.24	11.94	13.93	14.70	12.70
Al_2O_3	13.69	11.92	14.48	3.04	2.72	3.11	3.37
TiO ₂	5.16	3.12	2.57	11.34	11.29	11.86	11.25
FeO*	14.15	11.59	11.86	0.16	0.16	0.16	0.15
MnO	0.18	0.16	0.16	9.70	10.35	9.18	9.26
CaO	9.76	10.05	10.19	13.24	9.34	10.16	11.01
MgO	5.46	13.21	9.62	1.61	0.94	1.08	2.10
K ₂ O	1.07	0.48	0.96	2.69	2.73	2.62	3.28
Na ₂ O	3.35	3.10	2.64	0.44	0.29	0.31	0.56
P_2O_5	0.50	0.44	0.27	71.01	63.44	64.24	67.25
Mg#	44.72	70.51	62.98	11.94	13.93	14.70	12.70
Trace Elemen					0-2		
Ni	35	386	214	385	174	202	281
Cr	4	603	339	600	423	526	440
V	389	238	252	233	251	265	224
Ga	25	19	21	19	20	22	21
Cu	84	76	73	77	81	54	63
Zn	160	120	121	115	110	122	132
Cs	0.28	0.40	0.14	0.43	0.14	0.13	0.62
Rb	25.0	55.9	23.2	48.0	25.1	16.8	55.2
Ba	213	431	235	437	224	323	543
Th	2.80	3.36	2.18	3.48	2.41	2.23	4.93
U	0.82	0.70	0.43	0.81	0.57	0.73	1.19
Nb	40.56	42.85	29.16	44.36	28.81	36.51	67.26
Ta	2.59	2.65	1.80	2.70	1.76	2.58	4.15
La	30.24	31.95	41.87	31.95	22.82	20.37	41.90
Ce	72.09	69.29	47.58	69.49	49.31	44.97	87.77
Pb	3.15	2.79	2.01	2.87	2.07	2.06	4.22
Pr	9.91	8.76	8.94	8.80	6.36	5.94	10.91
Nd	42.30	34.84	36.05	34.74	25.53	24.21	41.94
Sr	552	585	436	821	419	394	679
Zr	301	188	161	191	156	189	267
Hf	7.55	4.71	4.00	4.69	3.90	4.74	6.35
Sm	9.98	7.32	7.55	7.29	5.70	5.50	8.42
Eu	3.23	2.35	2.67	2.33	1.92	1.83	2.62
Gd	9.57	6.67	9.17	6.60	5.62	5.38	7.48
Tb	1.37	0.92	1.32	0.92	0.83	0.81 4.64	1.03 5.47
Dy	7.65	4.99	7.93	4.93	4.74		0.95
Но	1.38	0.87	1.59	0.86	0.87	0.86	23.34
Y	33.23	20.89	46.87	20.84	21.40	21.05	2.24
Er	3.36	2.06	4.10	2.07	2.20	2.19	
Tm	0.44	0.27	0.54	0.26	0.29	0.30	0.28
Yb	2.50	1.49	3.09	1.49	1.71	1.72	1.59
Lu	0.35	0.21	0.49	0.21	0.25	0.25	0.22
Sc	26.0	22.2	24.4	21.4	25.2	21.5	18.7
86Sr/87Sr	0.705179	0.705823	0.706670	0.705856	0.706338		0.706528
43 Nd/144 Nd	0.512883	0.512702		0.512698	0.512678	0.512725	0.512682
²⁰⁶ Pb/ ²⁰⁴ Pb	18.848	18.801		18.804	18.866	18.821	18.803
²⁰⁷ Pb/ ²⁰⁴ Pb	15.564	15.609		15.612	15.613	15.604	15.615
10/ 10	13.304	15.005		10.012	15.015	10.004	

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Volcano	Savai'i PE	Savai'i PE	Savai'i PE	Savai'i PE	Savai'i PE	Savai'i PE	Savai'i Pl
Sample	S 15 Ma	S 26 S	S 28 Pb	S 31 P	S 32 M	S 33 P	S 34 A
Volcanic		10000					
Series	Mulifanua	Salani	Puapua	Puapua	Mulifanua	Puapua	Aopo
Lat. (°S)	13.3094	13.4327	13.4611	13.4753	13.3799	13.3099	13.318
Long (°W)	172.7021	172.314	172.3728	172.513	172.6655	172.7404	172.5523
Major eleme							
SiO ₂	44.62	45.51	48.64	48.76	45.39	48.05	45.89
Al_2O_3	12.40	12.64	14.20	14.18	11.84	14.25	12.38
TiO_2	3.30	3.24	3.01	2.96	3.83	2.73	3.13
FeO*	11.46	11.80	10.58	10.48	12.11	10.91	11.23
MnO	0.16	0.15	0.14	0.14	0.15	0.15	0.16
CaO	10.48	9.07	8.07	8.05	9.07	8.57	9.51
MgO	11.92	12.04	8.98	9.08	12.75	9.76	12.31
K_2O	1.91	1.80	2.09	2.06	1.88	1.68	1.78
Na ₂ O	3.04	3.25	3.73	3.72	2.62	3.28	3.15
P_2O_5	0.71	0.52	0.56	0.56	0.36	0.61	0.47
Mg#	68.57	68.14	64.03	64.50	68.82	65.24	69.68
Trace Elemen							
Ni	275	329		233	361	236	340
Cr	500	405		315	520	350	535
V	244	238		196	268	201	240
Ga	21	22		23	22	22	20
Cu	69	59		55	75	50	60
Zn	128	139		133	134	129	118
Cs	0.58	0.52		0.75	0.65	0.48	0.50
Rb	51.0	46.1		61.4	55.2	44.9	49.3
Ba	500	388		451	399	503	401
Th	6.34	3.39		5.33	2.82	4.75	3.94
U	1.43	0.89		1.39	0.73	1.11	0.87
Nb	70.27	54.93		55.33	50.96	55.28	50.26
Ta	4.27	3.42		3.55	3.63	3.43	3.28
La	54.90	32.20		36.37	24.26	39.62	36.60
Ce	112.66	71.92		75.05	54.51	81.56	77.21
Pb	5.24	3.06		4.21	2.32	3.94	3.08
Pr	13.50	9.34		9.26	7.17	9.82	9.48
Nd	50.24	37.53		35.63	29.73	37.02	36.57
Sr	748	670		610	491	606	556
Zr	272	251		284	250	253	225
Hf	6.39	6.00		6.64	6.40	5.93	5.57
Sm	9.59	8.09		7.55	6.66	7.50	7.58
Eu	2.92	2.57		2.43	2.15	2.37	2.30
Gd	8.11	7.31		7.00	6.18	6.83	6.88
Tb	1.11	1.02		0.99	0.86	0.98	0.95
Dy	5.78	5.35		5.31	4.60	5.46	5.15
Но	0.99	0.91		0.91	0.79	0.98	0.90
Y	24.41	21.99		22.24	19.21	24.02	21.52
Er	2.39	2.10		2.14	1.87	2.39	2.11
Tm	0.30	0.26		0.28	0.24	0.31	0.27
Yb	1.67	1.43		1.54	1.31	1.84	1.53
Lu	0.24	0.20		0.22	0.18	0.26	0.22
Sc	21.3	18.4		15.8	20.4	17.2	20.3
⁸⁶ Sr/ ⁸⁷ Sr ⁴³ Nd/ ¹⁴⁴ Nd ⁰⁶ Pb/ ²⁰⁴ Pb	0.706435	0.705597	0.705784	0.705765	0.705267	0.705451	0.706409
Nd/ Nd	10.000	10.702	0.512747	0.512751	0.512795	0.512776	
⁰⁷ Pb/ ²⁰⁴ Pb	18.888	18.793	18.776	18.777	18.842	18.682	18.796
⁰⁸ Pb/ ²⁰⁴ Pb	15.623 39.163	15.580	15.602	15.598	15.589	15.603	15.607
ru/ Pu	39.103	38.866	38.938	38.921	39.017	38.856	39.051

Volcano	Savai'i PE	Upolu PE					
Sample	S 35 P	S 39 A	S 42 A	S 43 P	S 50 S	S 52 M	U 10 S
Volcanic							
Series	Puapua	Aopo	Aopo	Puapua	Salani	Mulifanua	Salani
Lat. (°S)	13.2957	13.2703	13.2673	13.32	13.3596	13.42	13.8368
Long.(°W)	172.4654	172.3123	172.3106	172.2362	172.2673	172.2024	171.7362
Major eleme	nts (wt%)						
SiO ₂	48.30	47.33	46.05	44.72	45.94	45.83	48.41
Al_2O_3	13.32	12.47	11.76	11.45	12.59	11.98	15.15
TiO ₂	2.94	2.70	2.83	3.67	3.27	3.04	4.23
FeO*	11.29	11.71	11.98	12.42	12.37	11.45	11.88
MnO	0.15	0.16	0.16	0.16	0.17	0.15	0.16
CaO	9.44	9.45	9.28	9.13	9.79	9.24	9.64
MgO	10.16	12.04	13.75	14.06	11.95	13.33	5.21
K ₂ O	1.23	0.96	1.19	1.56	1.32	1.59	1.40
Na ₂ O	2.84	2.88	2.66	2.45	2.32	2.86	3.26
P_2O_5	0.33	0.31	0.35	0.39	0.27	0.52	0.65
Mg#	65.38	68.32	70.64	70.37	66.94	70.93	47.92
Trace Elemen							
Ni	255	340	412		302	402	62
Cr	456	570	594		552	491	86
V	236	231	231		266	223	309
Ga	21	20	19		21	20	25
Cu	55	72	66		83	65	78
Zn	114	121	124		123	126	146
Cs	0.15	0.30	0.36		0.51	0.66	0.31
Rb	29.5	24.8	31.0		38.6	44.0	33.7
Ba	277	245	288		314	438	298
Th	2.44	2.44	2.91		1.88	4.35	3.87
U	0.55	0.57	0.70		0.46	1.06	1.08
Nb	38.69	31.82	36.65		47.12	51.33	52.29
Ta	2.73	2.12	2.40		3.45	3.13	3.27
La	21.29	22.02	26.43		19.08	37.49	42.54
Ce	47.10	47.50	56.59		44.93	78.41	97.06
Pb	3.18	1.98	2.38		1.72	3.78	3.03
Pr	6.10	6.08	7.15		6.04	9.67	12.72
Nd	25.00	24.42	28.34		25.09	37.30	51.10
Sr	402	398	437		409	794	676
Zr	187	159	175		211	225	326
Hf	4.95	4.16	4.47		5.53	5.39	7.96
Sm	5.76	5.60	6.13		5.62	7.70	10.63
Eu	1.90	1.81	1.92		1.85	2.44	3.33
Gd	5.70	5.57	5.84		5.38	6.94	9.68
Tb	0.84	0.83	0.85		0.77	0.97	1.37
Dy	4.75	4.74	4.73		4.29	5.20	7.50
Но	0.86	0.87	0.85		0.77	0.90	1.35
Y	20.55	20.92	20.49		18.63	21.87	32.62
Er	2.12	2.16	2.09		1.90	2.15	3.32
Tm	0.28	0.28	0.27		0.25	0.27	0.44
Yb	1.60	1.61	1.54		1.43	1.53	2.50
Lu	0.23	0.23	0.22		0.20	0.21	0.35
Sc	21.5	22.1	20.8		22.5	19.0	21.6
86Sr/87Sr		0.705564	0.705793	0.706018	0.706907	0.705686	0.705363
43Nd/144Nd		0.512705	0.1.00170	0.512712	0.512711	0.512736	0.512870
206Pb/ ²⁰⁴ Pb	10.053		10.700	0.312/12			
	18.853	18.803	18.792		18.901	18.746	18.940
²⁰⁷ Pb/ ²⁰⁴ Pb	15.602	15.607	15.607		15.611	15.602	15.577
²⁰⁸ Pb/ ²⁰⁴ Pb	39.071	39.039	39.044		39.150	38.916	38.826

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Volcano	Upolu PE	Upolu PE	Upolu PE	Upolu PE	Upolu PE	Upolu PE	Upolu PE
Sample	U 15 S	U 16 S	U 17 S (1)	U 17 S (2)	U 21 S	U 23 P	U 24 L
Volcanic							
Series	Salani	Salani	Salani	Salani	Salani	Puapua	Lefaga
Lat. (°S)	13.9983	14.0306	14.0357	Same Flow	13.9541	14.0221	13.9668
Long. (°W)	171.4222	171.4325	171.4619	Few meters	171.7674	171.731	171.9434
Major elemen						Section Control	
SiO ₂	44.68	47.20	45.13		45.29	41.85	44.11
Al_2O_3	12.02	12.21	11.67		12.26	11.79	12.63
TiO ₂	4.53	3.99	3.96		3.45	4.52	3.62
FeO*	12.55	12.32	12.33		12.61	13.48	12.32
MnO	0.16	0.15	0.16		0.17	0.17	0.17
CaO	9.96	8.99	9.87		10.45	10.91	10.83
MgO	11.63	10.57	11.94		11.51	11.14	11.04
K ₂ O	1.66	1.54	1.51		1.19	1.85	1.40
Na ₂ O	2.21	2.67	2.88		2.52	3.56	3.15
P_2O_5	0.59	0.36	0.56		0.55	0.72	0.72
Mg#	66.02	64.28	67.00		65.68	63.42	65.27
Trace Elemen							
Ni	315		327		284	248	240
Cr	427		430		441	325	419
V	275		257		245	288	270
Ga	22		20		21	21	21
Cu	21		81		67	66	69
Zn	151		134		131	145	128
Cs	0.19		0.47		0.43	0.63	0.19
Rb	43.7		56.8		31.9	53.3	38.4
Ba	533		466		403	677	486
Th	4.41		3.73		4.54	6.73	7.17
U	1.06		0.87		1.03	1.43	0.93
Nb	50.54		45.98		45.85	72.56	54.02
Ta	3.09		2.85		2.80	4.09	3.03
La	41.63		36.96		42.94	63.55	65.36
Ce	88.37		79.28		89.24	129.64	130.00
Pb	2.26		2.99		3.68	4.95	4.48
Pr	11.01		9.96		11.00	15.49	14.99
Nd	43.34		39.79		42.36	57.75	53.99
Sr	643		613		623	892	823
Zr	219		203		225	264	231
Hf	5.49		5.24		5.70	6.36	5.40
Sm	9.01		8.54		8.54	11.03	10.00
Eu	2.76		2.61		2.60	3.37	3.00
Gd	8.27		7.84		7.52	9.40	8.39
Tb	1.13		1.09		1.04	1.28	1.16
Dy	6.06		5.89		5.69	6.70	6.23
Но	1.04		1.02		0.99	1.15	1.09
Y	24.89		24.32		23.80	27.84	27.05
Er	2.41		2.38		2.35	2.64	2.63
Tm	0.30		0.30		0.30	0.32	0.33
Yb	1.67		1.66		1.67	1.78	1.94
Lu	0.23		0.23		0.23	0.24	0.27
Sc	21.3		21.2		21.4	21.5	22.3
⁸⁶ Sr/ ⁸⁷ Sr		0.706088	0.705813	0.705750	0.705393	0.705536	0.705670
43Nd/144Nd			0.512711	0.512713	0.512662	0.512671	0.512666
206Pb/204Pb	10 705			0.512/15	0.512002		0.312000
	18.785		18.711			18.577	
²⁰⁷ Pb/ ²⁰⁴ Pb	15.623		15.607			15.611	
²⁰⁸ Pb/ ²⁰⁴ Pb	39.004		38.871			38.775	

Volcano	Upolu PE	Upolu PE	Upolu PE	Upolu PE	Upolu PE	Upolu PE	Upolu PE
Sample	U 26 L	U 28 L	U 30 L	U 32 M	U 33 M	U 35 M (1)	U 35 M (2
Volcanic							
Series	Lefaga	Lefaga	Lefaga	Mulifanua	Mulifanua	Mulifanua	Mulifanua
Lat. (°S)	13.9728	13.9677	13.9804	13.7917	13.8265	13.8359	Same Flow
Long.(°W)	171.9399	171.9416	171.9329	171.9007	171.9974	171.9469	Few meter
Major eleme	nts (wt%)						
SiO ₂	41.64	42.25	42.10	44.65	48.37	41.58	
Al_2O_3	11.95	11.58	11.74	12.14	13.63	11.14	
TiO ₂	5.82	5.63	5.67	4.31	3.34	6.58	
FeO*	14.39	13.92	14.02	13.14	11.30	14.55	
MnO	0.18	0.16	0.17	0.16	0.14	0.17	
CaO	10.26	9.90	10.03	9.67	9.01	9.97	
MgO	11.67	11.44	11.45	12.35	8.12	11.77	
K_2O	1.20	1.65	1.54	1.27	1.92	1.39	
Na ₂ O	2.29	2.87	2.69	1.99	3.50	2.42	
P_2O_5	0.61	0.59	0.58	0.34	0.68	0.42	
Mg#	62.99	63.29	63.14	66.35	60.11	62.92	
Trace Eleme				100			
Ni	299	301		357	171	328	
Cr	376	375		489	204	460	
V	322	320		292	189	379	
Ga	22	22		20	23	20	
Cu	67	69		68	29	87	
Zn	145	145		129	142	140	
Cs	0.54	0.54		0.29	0.51	0.41	
Rb	37.8	50.2		32.3	47.9	40.4	
Ba	509	499		425	576	429	
Th	5.00	4.90		2.26	4.86	3.81	
U	1.07	1.08		0.47	1.14	0.84	
Nb	61.34	60.31		35.56	49.91	48.46	
Та	3.73	3.66		2.40	2.84	3.08	
La	47.75	47.09		24.93	44.65	36.18	
Ce	100.15	98.79		52.55	91.89	77.47	
РЬ	3.51	3.37		1.94	4.03	2.74	
Pr	12.26	12.15		7.00	11.32	9.72	
Nd	46.86	46.38		28.67	44.05	37.78	
Sr	704	689		641	766	524	
Zr	260	256		186	235	226	
Hf	6.51	6.33		4.99	5.72	6.02	
Sm	9.58	9.39		6.48	9.30	7.92	
Eu	2.95	2.88		2.08	2.98	2.46	
Gd	8.50	8.32		6.25	8.54	7.21	
Tb	1.18	1.15		0.89	1.19	1.01	
Dy	6.19	6.08		4.83	6.27	5.56	
Но	1.08	1.06		0.87	1.07	0.99	
Y	25.96	25.48		21.69	26.33	23.61	
Er	2.49	2.44		2.09	2.46	2.35	
Tm	0.31	0.30		0.27	0.30	0.30	
Yb	1.73	1.68		1.53	1.64	1.74	
Lu	0.24	0.23		0.21	0.22	0.23	
Sc	22.8	22.6		23.8	16.1	27.4	
86Sr/87Sr	0.705620	0.705594	0.705379	0.706113	0.705664	0.705631	0.705607
43Nd/144Nd	0.512728	0.512684	0.512881	0.512734	0.512690	0.512668	0.512656
²⁰⁶ Pb/ ²⁰⁴ Pb		18.696				18.681	
²⁰⁷ Pb/ ²⁰⁴ Pb							
		15.610				15.606	
²⁰⁸ Pb/ ²⁰⁴ Pb		38.878				38.883	

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<i>Table 2-5</i> ,				
Volcano	Upolu PE	I	BHVO-1 Standar	rd
Sample	U 37 P			
Volcanic				
Series	Puapua			
Lat. (°S)	14.0068			
Long. (°W)	171.7292	Long-ter	m average, U. Q	ueensland
Major elemen				
SiO_2	41.75			
Al_2O_3	11.84			
TiO ₂	4.45	Ti 49	16376	1.02
FeO*	13.34			
MnO	0.18			
CaO	10.92			
MgO	11.06			
K_2O	1.86			
Na ₂ O	3.92			
P_2O_5	0.69			
Mg#	63.49			
Trace Elemen			11.00	2
Ni	241	Ni 60	116	2.57
Cr	312	Cr 52	295	2.37
V	290	V 51	286	1.39
Ga	21	Ga 71	21.2	1.18
Cu	67	Cu 65	137	0.94
Zn	142	Zn 66	106	1.81
Cs	0.63	Cs 133	0.0967	1.21
Rb	53.6	Rb 85	9.27	0.83
Ba	678	Ba 137	132	0.75
Th	6.58	Th 232	1.19	1.09
U Nb	1.39	U 238	0.433	1.09
Ta	71.19	Nb 93	18.4	0.86
La	4.04	Ta 181	1.15	1.31
Ce	61.31 125.23	La 139	15.1	0.74
Pb	95.09	Ce 140 Pb 208	37.7 1.97	0.65
Pr	15.00	Pr 141	5.45	3.87 0.55
Nd	55.86	Nd 146	24.1	0.52
Sr	861	Sr 86	394	0.52
Zr	256	Zr 90	165	0.94
Hf	6.20	Hf 178	4.37	1.28
Sm	10.69	Sm 149	5.95	0.79
Eu	3.27	Eu 151	2.04	0.98
Gd	9.10	Gd 160	5.97	0.68
Tb	1.25	Tb 159	0.886	0.97
Dy	6.59	Dy 161	5.11	0.85
Но	1.13	Ho 165	0.953	0.84
Y	27.60	Y 89	22.9	0.76
Er	2.63	Er 167	2.42	1.32
Tm	0.32	Tm 169	0.324	1.28
Yb	1.79	Yb 172	1.90	1.05
Lu	0.24	Lu 175	0.268	1.36
Sc	22.2	Sc 45	29.9	1.68
⁸⁶ Sr/ ⁸⁷ Sr	0.705492			
143Nd/144Nd	0.512618			
²⁰⁶ Pb/ ²⁰⁴ Pb	0.512010			
²⁰⁷ Pb/ ²⁰⁴ Pb				
²⁰⁸ Pb/ ²⁰⁴ Pb				

Table 2-6. Isotopic compositions of samples previously collected by KAF**.

91TP-128 upper shield 0.705355 0.512846 -	Volcano	Sample	Type	$^{86}\mathrm{Sr}/^{87}\mathrm{Sr}$	$^{143}Nd/^{144}Nd$	³ He/ ⁴ He	$^{206}\mathrm{Pb}/^{204}\mathrm{Pb}$	$^{206}Pb/^{204}Pb ^{207}Pb/^{204}Pb ^{208}Pb/^{204}Pb$	$^{208}\mathrm{Pb}/^{204}\mathrm{Pb}$
91TP-134 dike 0.705195 0.512821 24.67 19.065 15.614 91TP-165 dike 0.705166 0.5128 25.79 - - 91TP-201 dike 0.706491 0.51285 25.79 - - 91TP-203 lower shield 0.706493 0.512895 14.44 - - 91TP-203 lower shield 0.707143 0.512895 14.83 19.106 15.625 91TP-204 upper shield 0.707476 0.512898 - - - 91TP-196 upper shield 0.706374 0.512898 - - - 91TP-202 upper shield 0.705374 0.51283 - - - 91TP-204 upper shield 0.706354 0.51283 - - - - 91TP-252 upper shield 0.706354 0.51283 - - - - 91TP-252 upper shield 0.706359 0.51283 - -	Tutuila	91TP-128	upper shield	0.705535	0.512846		1	ï	1
91TP-165 dike 0.705166 0.5128 25.79 - 91TP-201 dike 0.704971 0.512825 20.49 19.199 15.59 91TP-201 lower shield 0.704691 0.512825 14.84 19.106 15.525 91TP-103 lower shield 0.707863 0.512844 13.97 - - 91TP-104 upper shield 0.704476 0.512848 - - - 91TP-105 upper shield 0.705317 0.512863 - - - 91TP-207 upper shield 0.706324 0.51286 - - - - 91TP-208 upper shield 0.706349 0.512863 1.2.44 - - - 91UF-80 shield 0.706349 0.512783 11.44 - - - 91UF-80 shield 0.706349 0.512783 11.27 - - 91UF-80 shield 0.706349 0.51285 1.2.28 19.04<	Tutuila	91TP-134	dike	0.705195	0.512821	24.67	19.065	15.614	39.13
91TP-201 dike 0.704971 0.512825 20.49 19.199 15.59 91TP-133 lower shield 0.70609 0.512709 17.44 - - 91TP-203 lower shield 0.707143 0.512858 - - - 91TP-203 lower shield 0.707317 0.512858 - - - 91TP-144 upper shield 0.705317 0.512858 - - - 91TP-156 upper shield 0.70525 0.512453 - - - 91TP-228 upper shield 0.706324 0.51287 - - - 91TP-252 upper shield 0.706349 0.512657 14.44 - - 91TP-26 upper shield 0.706349 0.512783 11.44 - - 91UF-86 shield 0.706349 0.512783 11.44 - - 91UF-87 shield 0.706349 0.512783 11.27 - -	Tutuila	91TP-165	dike	0.705166	0.5128	25.79		ı	
91TP-133 lower shield 0.706609 0.512709 17.44 - - 91TP-203 lower shield 0.707863 0.512595 14.83 19.106 15.625 91TP-207 lower shield 0.707317 0.51284 13.97 - - 91TP-144 upper shield 0.70476 0.51283 - - - 91TP-228 upper shield 0.70525 0.512453 - - - 91TP-228 upper shield 0.70634 0.512657 14.44 - - 91TP-529 upper shield 0.70634 0.51283 12.59 19.124 15.645 91TP-63B shield 0.70634 0.512783 11.4 18.974 15.615 91UF-63B shield 0.706345 0.512851 16.7 - - 91UF-64B shield 0.706447 0.51285 12.28 18.988 15.627 91UF-65B shield 0.706447 0.51283 12.28 18.988	Tutuila	91TP-201	dike	0.704971	0.512825	20.49	19.199	15.59	39.171
91TP-203 lower shield 0.707863 0.512595 14.83 19.106 15.625 91TP-207 lower shield 0.707143 0.512644 13.97 - - 91TP-144 upper shield 0.705317 0.512848 - - - 91TP-144 upper shield 0.706476 0.512838 - - - 91TP-252 upper shield 0.706324 0.512657 - - - 91TP-252 upper shield 0.706349 0.512863 - - - 91TP-254 upper shield 0.706347 0.512783 11.44 - - 91UF-65B shield 0.70647 0.512725 10.25 - - 91UF-86 shield 0.70547 0.512851 16.7 - - 91UF-89 shield 0.70547 0.512851 16.7 - - 91UF-89 shield 0.70547 0.51285 12.28 18.98 15.627 <tr< th=""><th>Tutuila</th><th>91TP-133</th><th>lower shield</th><th>0.70609</th><th>0.512709</th><th>17.44</th><th></th><th>ı</th><th>,</th></tr<>	Tutuila	91TP-133	lower shield	0.70609	0.512709	17.44		ı	,
91TP-207 lower shield 0.707143 0.512644 13.97 - - 91TP-144 upper shield 0.705317 0.512858 - - - 91TP-144 upper shield 0.705225 0.512453 - - - 91TP-252 upper shield 0.70534 0.512657 14.44 - - 91TP-252 upper shield 0.70534 0.512657 14.44 - - 91TP-252 upper shield 0.70534 0.512803 22.59 19.124 15.615 91UF-80B shield 0.705349 0.512725 10.25 - - 91UF-80B shield 0.705245 0.512725 10.25 - - 91UF-80B shield 0.705447 0.512851 16.78 - - 91UF-80B shield 0.706447 0.51283 11.27 - - 91UF-80B shield 0.706447 0.512833 15.23 18.978 15.627 <th>Tutuila</th> <th>91TP-203</th> <th>lower shield</th> <th>0.707863</th> <th>0.512595</th> <th>14.83</th> <th>19.106</th> <th>15.625</th> <th>39.526</th>	Tutuila	91TP-203	lower shield	0.707863	0.512595	14.83	19.106	15.625	39.526
91TP-144 upper shield 0.705317 0.512858 -	Tutuila	91TP-207	lower shield	0.707143	0.512644	13.97	1	1	1
91TP-196 upper shield 0.704476 0.5129 - 18.806 15.543 91TP-228 upper shield 0.705225 0.512453 - - - 91TP-252 upper shield 0.706354 0.512657 14.44 - - 91TPK-5 upper shield 0.706392 0.512863 12.59 19.124 15.615 91UF-86 shield 0.706389 0.512725 10.25 - - 91UF-86 shield 0.706389 0.512725 10.25 - - 91UF-89 shield 0.706349 0.512851 16.78 - - 91UF-80 shield 0.706447 0.512851 16.78 - - 91UF-84 shield 0.706447 0.512851 15.28 18.988 15.627 UPO-10C shield 0.706431 0.512833 15.8 - - UPO-F-19 shield 0.70493 0.512833 15.29 - - <tr< th=""><th>Tutuila</th><th>91TP-144</th><th>upper shield</th><th>0.705317</th><th>0.512858</th><th>,</th><th></th><th>ı</th><th>ı</th></tr<>	Tutuila	91TP-144	upper shield	0.705317	0.512858	,		ı	ı
91TP-228 upper shield 0.705225 0.512453 - - - 91TP-252 upper shield 0.706354 0.512657 14.44 - - 91TPK-5 upper shield 0.705192 0.512803 22.59 19.124 15.646 91UF-65B shield 0.706349 0.512725 10.25 - - 91UF-86 shield 0.706389 0.512725 10.25 - - 91UFK-4 shield 0.70545 0.512851 16.78 - - 91UFK-6A shield 0.705147 0.512851 16.78 - - 91UFK-6A shield 0.70647 0.512851 11.27 - - 91UFK-6A shield 0.70647 0.512853 11.27 - - 91UFK-6A shield 0.70647 0.512833 15.8 - - 0 UPO-F-19 shield 0.70649 0.512833 15.29 - - 0 UP	Tutuila	91TP-196	upper shield	0.704476	0.5129	r	18.806	15.543	38.656
91TP-252 upper shield 0.706354 0.512657 14.44 - - 91TPK-5 upper shield 0.705192 0.512803 22.59 19.124 15.646 91UF-65B shield 0.70547 0.512783 11.4 18.974 15.615 91UF-86 shield 0.706389 0.512725 10.25 - - 91UFK-4 shield 0.70545 0.512851 16.7 - - 91UFK-6A shield 0.705147 0.512851 16.7 - - 91UFK-6A shield 0.706447 0.51285 12.28 18.988 15.627 UPO-1C shield 0.706447 0.51283 15.8 - - UPO-F-19 shield 0.706449 0.51283 15.29 - - UPO-F-20 shield 0.704569 0.51283 15.23 - - 82MT11 shield 0.704669 0.512783 13.34 - -	Tutuila	91TP-228	upper shield	0.705225	0.512453				1
91TPK-5 upper shield 0.705192 0.512803 22.59 19.124 15.646 91UF-65B shield 0.706347 0.512783 11.4 18.974 15.615 91UF-86 shield 0.705245 0.512725 10.25 - - 91UF-89B shield 0.705147 0.512851 16.78 - - 91UFK-4 shield 0.705147 0.512851 16.7 - - 91UFK-6A shield 0.706447 0.51285 12.28 18.988 15.627 UPO-10C shield 0.706447 0.51283 15.8 - - UPO-F-19 shield 0.706447 0.512833 15.8 - - UPO-F-20 shield 0.70453 0.512859 12.92 - - UPO-F9-7 shield 0.704569 0.512833 15.23 - - 82MT11 shield 0.704661 0.512783 13.34 - - <t< th=""><th>Tutuila</th><th>91TP-252</th><th>upper shield</th><th>0.706354</th><th>0.512657</th><th>14.44</th><th>1</th><th>ī</th><th>ı</th></t<>	Tutuila	91TP-252	upper shield	0.706354	0.512657	14.44	1	ī	ı
91UF-65B shield 0.70547 0.512783 11.4 18.974 15.615 91UF-86 shield 0.706389 0.512725 10.25 - - 91UF-89B shield 0.705245 0.512772 16.78 - - 91UFK-4 shield 0.705147 0.512851 16.7 - - 91UFK-6A shield 0.706447 0.51285 12.28 18.988 15.627 UPO-10C shield 0.70522 0.51283 15.8 - - UPO-F-19 shield 0.70522 0.512833 15.8 - - UPO-F-20 shield 0.704569 0.512833 15.29 - - UPO-F-20 shield 0.704649 0.512783 13.34 - - 82MT11 shield 0.704649 0.512783 13.34 - - 82MT8B shield 0.704649 0.512776 19.06 19.316 - 91SVK7	Tutuila	91TPK-5	upper shield	0.705192	0.512803	22.59	19.124	15.646	39.224
91UF-86 shield 0.705345 0.512725 10.25 - - 91UF-89B shield 0.705245 0.512772 16.78 - - 91UFK-4 shield 0.705147 0.512851 16.7 - - 91UFK-6A shield 0.704821 0.51285 12.28 18.988 15.627 UPO-F-19 shield 0.706447 0.512833 15.28 - - UPO-F-20 shield 0.704593 0.512833 15.8 - - UPO-F-7 shield 0.704569 0.512833 15.23 - - 82MT11 shield 0.704649 0.512783 15.23 - - 82MT17 shield 0.704649 0.512783 13.34 - - 82MT18 shield 0.704649 0.512781 13.54 - - 82MT8B shield 0.704637 0.51277 - - - 91SVK7 COBBLE	Upolu	91UF-65B	shield	0.70547	0.512783	11.4	18.974	15.615	39.126
91UF-89B shield 0.705245 0.512772 16.78 - - 91UFK-4 shield 0.704821 0.512851 16.7 - - 91UFK-6A shield 0.704821 0.51285 12.28 18.988 15.627 UPO-10C shield 0.706447 0.512845 11.27 - - UPO-F-19 shield 0.70522 0.512859 12.92 - - UPO-F-2 shield 0.704569 0.512859 12.92 - - 82MT11 shield 0.704649 0.512833 15.23 - - 82MT17 shield 0.704649 0.512783 13.34 - - 82MT18 shield 0.704641 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.51277 - - - 91SVK7 COBBLE 0.7072 0.51277 - 18.872 15.587	Upolu	91UF-86	shield	0.706389	0.512725	10.25	1	,	
91UFK-4 shield 0.705147 0.512851 16.7 - - 91UFK-6A shield 0.704821 0.51285 12.28 18.988 15.627 UPO-10C shield 0.706447 0.512743 11.27 - - UPO-F-19 shield 0.70522 0.512833 15.8 - - UPO-F-20 shield 0.70493 0.512859 12.92 - - UPO-F9-7 shield 0.704569 0.512833 15.23 - - 82MT11 shield 0.704649 0.512783 13.34 - - 82MT18 shield 0.704649 0.512783 13.54 - - 82MT18 shield 0.704661 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.51277 - 18.872 15.587	Upolu	91UF-89B	shield	0.705245	0.512772	16.78		i	1
91UFK-6A shield 0.704821 0.51285 12.28 18.988 15.627 UPO-10C shield 0.706447 0.512743 11.27 - - UPO-F-19 shield 0.70522 0.512833 15.8 - - UPO-F-20 shield 0.70493 0.51281 12.92 - - UPO-F9-7 shield 0.704649 0.512833 15.23 - - 82MT11 shield 0.704649 0.512783 13.34 - - 82MT18 shield 0.704661 0.512781 13.54 - - 82MT8B shield 0.704661 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.512771 - - - 91SVK7 COBBLE 0.7072 0.51277 - 18.872 15.587	Upolu	91UFK-4	shield	0.705147	0.512851	16.7	1		
UPO-10C shield 0.706447 0.512743 11.27 - - UPO-F-19 shield 0.705133 0.512833 15.8 - - UPO-F-19 shield 0.70522 0.512859 12.92 - - UPO-F-20 shield 0.70493 0.512912 13.8 - - 82MT11 shield 0.704649 0.512833 15.23 - - 82MT18 shield 0.704649 0.512781 13.34 - - 82MT18 shield 0.704661 0.512781 13.54 - - 82MT8B shield 0.704637 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.512771 - 18.872 15.587	Upolu	91UFK-6A	shield	0.704821	0.51285	12.28	18.988	15.627	39.171
UPO-F-19 shield 0.705133 0.512833 15.8 - - UPO-F-20 shield 0.70493 0.512859 12.92 - - UPO-F-20 shield 0.704693 0.512912 13.8 - - SZMT11 shield 0.704649 0.512783 15.23 - - SZMT18 shield 0.704649 0.512783 13.34 - - SZMT18 shield 0.704661 0.512781 13.54 - - SZMT8B shield 0.704637 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.512771 - 18.872 15.587	Upolu	UPO-10C	shield	0.706447	0.512743	11.27		,	
UPO-F-20 shield 0.70522 0.512859 12.92 - - UPO-F9-7 shield 0.704569 0.512833 15.23 - - 82MT11 shield 0.704649 0.512783 15.23 - - 82MT18 shield 0.704661 0.512781 13.34 - - 82MT8B shield 0.704661 0.512781 13.54 - - 91SVK-1 PE 0.704637 0.512776 19.06 19.316 15.593 91SVK7 COBBLE 0.7072 0.51277 - 18.872 15.587	Upolu	UPO-F-19	shield	0.705133	0.512833	15.8	r		
UPO-F9-7 shield 0.70493 0.512912 13.8 - - 82MT11 shield 0.704649 0.512833 15.23 - - 82MT17 shield 0.704649 0.512781 13.34 - - 82MT18 shield 0.704661 0.512781 13.54 - - 82MT8B shield 0.704637 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.512721 - - - 91SVK7 COBBLE 0.7072 0.51277 - 18.872 15.587	Upolu	UPO-F-20	shield	0.70522	0.512859	12.92	1	1	1
82MT11 shield 0.704569 0.512833 15.23	Upolu	UPO-F9-7	shield	0.70493	0.512912	13.8	1		1
82MT17 shield 0.704649 0.512783 13.34	Manua	82MT11	shield	0.704569	0.512833	15.23	,	1	1
82MT18 shield 0.704661 0.512781 13.54	Manua	82MT17	shield	0.704649	0.512783	13.34	•	•	
82MT8B shield 0.704637 0.512776 19.06 19.316 15.593 91SVK-1 PE 0.705589 0.512721 13.7	Manua	82MT18	shield	0.704661	0.512781	13.54	.1	,	r
91SVK-1 PE 0.705589 0.512721 13.7 18.872 15.587	Manua	82MT8B	shield	0.704637	0.512776	19.06	19.316	15.593	39.445
91SVK7 COBBLE 0.7072 0.51277 - 18.872 15.587	Savaii	91SVK-1	PE	0.705589	0.512721	13.7	1	1	
	Savaii	91SVK7	COBBLE	0.7072	0.51277		18.872	15.587	39.021

^{**} Analytical techniques as described in Farley et al. (1992).

Table 2-7. Calculated trace element composition of a "pure" EM2 melt.

	Average	Average	EM2
	Ta'u	Malumalu	Melt
⁸⁷ Sr/ ⁸⁶ Sr	0.7046	0.70889	0.7128
Cs	0.11	0.41	0.70
Rb	15.17	44.73	73.47
Ba	146.96	281.99	413.29
Th	2.78	6.17	9.46
U	0.62	1.16	1.69
Nb	29.71	43.96	57.83
Ta	2.06	2.92	3.76
K	0.73	1.57	2.39
La	24.16	44.31	63.90
Ce	50.61	82.17	112.85
Pb	2.09	4.35	6.56
Pr	6.27	9.16	11.98
Nd	27.24	34.89	42.33
Sr	367.00	379.09	390.83
Zr	187.15	205.43	223.20
Hf	4.82	5.20	5.57
Sm	6.74	7.15	7.56
Eu	2.15	1.99	1.84
Ti	2.99	2.20	1.42
Gd	6.41	6.28	6.15
Tb	0.95	0.87	0.79
Dy	5.21	4.59	4.00
Ho	0.94	0.80	0.67
Y	24.00	21.89	19.84
Er	2.26	1.91	1.57
Tm	0.29	0.25	0.21
Yb	1.67	1.40	1.13
Lu	0.24	0.21	0.17
Sc	25.76	25.77	25.79

^{**}All reported as ppm except K and Ti in wt%.

**All samples in averages are olivine corrected to Mg# 73.

Table 2-8. Source composition of EM2

		Depleted mantle			
	Primitive Mantle*	(2% melt depleted)	Bulk D values	EM2 Source	PUM
	(mdd)	(mdd)		(mdd)	Normalized
Rb	9.0	0.140	0.00001	1.456	2.427
Ba	9.9	1.587	0.00008	15.857	2.403
Th	0.0795	0.020	0.00058	0.177	2.224
n	0.0203	900'0	0.0016	0.040	1.968
NP	0.658	0.289	0.0040	1.087	1.653
Та	0.037	0.019	0.0042	0.063	1.698
La	0.648	0.287	0.0067	0.895	1.381
Ce	1.675	0.823	0.0117	1.923	1.148
Pb	0.15	0.027	0.0092	0.144	0.959
Pr	0.254	0.146	0.0214	0.251	0.987
PN	1.25	0.772	0.0317	1.140	0.912
Sr	19.9	10.803	0.0185	20.044	1.007
Zr	10.5	989.9	0.0472	8.835	0.841
H	0.283	0.195	0.0644	0.238	0.841
Sm	0.406	0.290	0.0692	0.347	0.855
Eu	0.154	0.114	0.1088	0.128	0.829
Ξ	1072.1	829.471	0.1428	900.5	0.840
PS	0.544	0.417	0.1800	0.445	0.818
Tb	0.099	0.079	0.2730	0.082	0.826
Dy	0.674	0.554	0.3504	0.569	0.845
Ho	0.149	0.125	0.4649	0.127	0.853
>	4.3	3.651	1.0829	3.655	0.850
Er	0.438	0.374	0.5708	0.378	0.864
Vb	0.441	0.387	1.0784	0.387	0.878

^{*} Primtive Upper Mantle (PUM) from McDonough and Sun (1995).

Chapter 3:

Major and Trace Element Composition of the Depleted MORB Mantle (DMM)*

Abstract

We derive an estimate for the chemical composition of the depleted MORB mantle (DMM), the source reservoir to mid-ocean ridge basalts (MORBs), which represents at least 30% the mass of the whole silicate Earth. A database for the chemical and physical properties of abyssal peridotites has become robust and complete enough to truly access a reference DMM. Using trace element depletion trends from the abyssal peridotites, it is possible to construct a large part of DMM's trace element pattern. Splicing this information with isotopic constraints (Sr-Nd-Pb-Hf) and canonical ratios (Ce/Pb, Nb/Ta, Nb/U, Ba/Rb, H₂O/Ce, CO₂/Nb and Cl/K), we can extend abundance estimates to all the incompatible elements including volatile content. The resulting trace element pattern for average DMM constrains parental MORB to be generated by 6% aggregated fractional melting – consistent with recent models for hydrous melting of the mantle (Asimow et al., 2004). We show that DMM is roughly balanced by the continental crust and better balanced upon inclusion of ocean island basalt source and oceanic crust components. Compared to the primitive mantle, DMM has been depleted by 2-3% melt extraction and has only 15% the radiogenic heat production.

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1. Introduction

The trace element composition of the depleted upper mantle (DMM) is a fundamental parameter in modeling the generation of mid-ocean ridge basalts (MORBs), calculating the crust-mantle mass balance, and establishing the chemical and thermal evolution of the Earth. To date, all attempts to establish an average composition for the upper mantle, whether by parent/daughter derivations from isotopic systems (Allègre et al., 1983; Galer and O'Nions; 1985; Albarede and Brouxel, 1987; Shimizu, 1998; Elliot et al., 1999; Salters and Stracke, 2004) or by MORB trace element inversions (Salters and Stracke, 2004; McKenzie and O'Nions, 1991, Hofmann, 1988, Sun and McDonough, 1989), have centered on MORB liquids and include major assumptions about melt generation, melt transport and differentiation processes that have affected these upper mantle melts. Unfortunately, it is just these processes that we are trying to understand, rather than assume *a priori*.

The most compelling evidence for upper mantle depletion comes from the heavy-element isotopic composition of MORBs. Although MORBs and their residues, abyssal peridotites, have some degree of heterogeneity in radiogenic isotope ratios (Sr-Nd-Pb-Hf), they are focused within a small range of values relative to ocean island basalts and are, with very few exceptions, depleted from bulk earth values in ⁸⁷Sr/⁸⁶Sr, ¹⁴³Nd/¹⁴⁴Nd and ¹⁷⁶Hf/¹⁷⁷Hf, thus requiring a long-term history of low Rb/Sr, Hf/Lu and Nd/Sm (i.e. incompatible element depletion). Direct evidence for upper mantle depletion came with the classic trace element studies of Johnson et al. (1990) and Johnson and Dick (1992) on abyssal peridotites, but most of the observed depletion in these peridotites is due to melt extraction during the latest spreading events, so that the composition of the general ambient upper mantle has been severely overprinted by this latest melting episode.

Ultimately, we know that this mantle, which has been estimated to comprise 30-70% by mass of the bulk silicate earth, has been depleted over time in the highly incompatible (lithophile) elements (Allègre et al., 1983; Hofmann, 1988, refs. therein). Most models ascribe the depletion of the upper mantle to the extraction of the enriched continental crust, a process that has removed up to 90% of the most incompatible elements, and 80-85% of the heat-producing elements (e.g. Hofmann, 1988). Some part of the depletion may also be created by the preservation of recycled oceanic crust that is currently sequestered in unknown regions of the mantle. It is possible the lower mantle (below 670 km) has also been depleted by such processes, but this question is highly controversial at present.

In place of assuming a set of bulk partition coefficients or a degree of melt extraction, we use in this paper the following constraints in deriving an average trace element composition of DMM: 1) trace element content of clinopyroxenes from abyssal peridotites, 2) isotopic evolution from primitive upper mantle (PUM) and 3) canonical trace element ratios in MORBs. Abyssal peridotite data from the literature, the backbone of this study, are treated in a way reminiscent of studies such as Loubet et al. (1975), Hart and Zindler (1986) and McDonough and Sun (1995). Abyssal peridotites, unlike basalts, are not modified by secondary processes such as fractional crystallization, magma mixing and crustal assimilation (Dick et al., 1984), but may be subject to melt impregnation (Dick, 1989) and melt/rock reaction (Kelemen et al., 1992). We show that melt impregnation can be recognized and therefore filtered from the abyssal peridotite dataset and suggest that melt/rock reaction in the dataset we have used has had more effect on major element chemistry than trace element chemistry. Overall, it seems that, for the moderately incompatible elements, abyssal peridotites more accurately record the trend of upper mantle depletion than do MORBs.

The average trace element content of DMM, as deduced here, generally shows a very smooth pattern with increasing trace element compatibility, which to first order mass-balances with the continental crust. The degree of depletion indicated in DMM represents 2-3% melt removal from the primitive upper mantle (PUM) of McDonough and Sun (1995); this means that DMM has only 15% the radiogenic heat production of PUM (from K, U and Th). Present-day ocean crust (i.e. MORBs) can be modeled with 6% aggregated fractional melting of the deduced DMM.

2. Trace Element Composition of DMM

2.1 Abyssal Peridotites

Abyssal peridotites have been shown to be residues of fractional or near-fractional melting that produces MORB (e.g. Johnson et al., 1990). Although bulk trace element compositions of these peridotites are heavily modified by alteration on the seafloor, many workers have analyzed the trace element compositions (Sr, Zr, Ti, Ce, Nd, Sm, Eu, Gd, Dy, Er, Yb, and Y) of relict, unaltered clinopyroxene grains as a means of chemical characterization (Johnson et al., 1990; Johnson and Dick, 1992; Dick and Natland, 1996; Hellebrand et al., 2002; Salters and Dick, 2002; Tartarotti et al., 2002). Using such

clinopyroxene compositions, bulk compositions ($C_{WholeRock}$) of the peridotites can be calculated for each trace element with the following equation:

$$C_{WholeRock} = C_{cpx} \left(\frac{D_{bulk}}{D_{cpx}} \right) \tag{1}$$

where the bulk partition coefficient (D_{bulk}) is determined from modal abundances ($x_{mineral}$) of olivine (ol), orthopyroxene (opx), clinopyroxene (cpx) and spinel (sp), and mineral/melt partition coefficients ($D_{mineral}$) as follows:

$$D_{bulk} = x_{ol} D_{ol} + x_{opx} D_{opx} + x_{cpx} D_{cpx} + x_{sp} D_{sp}$$
 (2)

This method of calculating bulk peridotite compositions is a substitute for a more accurate method, which would be to sum the concentrations of trace elements in minerals according to their modal proportions. Converting to whole rock compositions from clinopyroxene alone is done simply due to the fact that almost no trace element data exists for other minerals in the peridotite samples.

We have compiled a data set from the literature that includes abyssal peridotites having both cpx trace element analyses and modal abundances (data sources are Johnson et al., 1990; Johnson and Dick, 1992; Dick et al., 1984; Dick and Natland, 1996; Hellebrand et al., 2002; Salters and Dick, 2002; Tartarotti et al., 2002). All samples containing more than 1% modal plagioclase are excluded on the basis that most plagioclase in abyssal peridotites has been interpreted as a secondary phase representative of melt impregnation (Dick, 1989; Tartarotti et al., 2002); as a result, only 5 out of almost 90 samples included have any reported modal plagioclase at all. Anomalously high Ce and Sr values have also been excluded for some samples. For Sr, some anomalous values are suspected seawater alteration; such Sr anomalies are not accompanied by Eu anomalies and therefore are not suspected to be plagioclase reaction products. For both Sr and Ce, the most incompatible elements of the sample suite and therefore the most depleted in abyssal peridotites, elevated concentrations are probably due to small amounts of melt impregnation (see discussion below).

The final trace element data set for the abyssal peridotites, corrected to whole rock compositions using mineral/melt partition coefficients from Kelemen et al. (2003),

published modal abundances and the above equations, is shown in Figure 1 and the Supp. Data Table 1. It is important to note that the geographical distribution of the sample suite is highly focused on the Southern Ocean; 90% of the samples originate from the American-Antarctic, Southwest Indian and Central Indian Mid-Ocean Ridges, with the remaining 10% obtained from the Mid-Atlantic Ridge and East Pacific Rise. The question of whether or not this geographical bias manifests as a chemical bias will not be answered until larger datasets from other localities are established.

The purpose of compiling bulk rock information is to derive the inter-relationships between different trace elements (i.e. to what degree the various incompatible trace elements are depleted relative to each other over the entire history of mantle depletion). A formulation of the trace element "system" begins by interpreting the trace element compositions of peridotites using the equation for a residue of fractional melting (see Zou, 1998, for review of melting models). Calculations show that modal fractional melting used to interpret residues of melting is a fine approximation for any number of more complicated melting models (see Appendix 1). The ratio of the concentration of a given element in a solid, C_s , to its original concentration, C_o , after some fraction of melting, F_o , is as follows:

$$\frac{C_s}{C_a} = (1 - F)^{(\frac{1}{D} - 1)} \tag{3}$$

where D is the bulk solid/melt partition coefficient for modal melting.

The relationship between the concentrations of two elements in the residue, [A] and [B], can be linearized upon equating two equations solved for F:

$$\ln(C_s^A) = R \ln(C_s^B) + \ln \left(\frac{C_o^A}{\left(C_o^B\right)^R}\right)$$
(4)

where the slope, R, on the linear ln([A]) versus ln([B]) array is a function of the bulk partition coefficients for elements A and B:

$$R = \frac{D_B (1 - D_A)}{D_A (1 - D_B)} \tag{5}$$

Given that we have information for 12 different elements (see fig. 1), there are 132 of the ln([A]) vs. ln([B]) relationships to consider in the overall reduction of this dataset;

Figure 2 shows a sampling of these linear arrays, with the more incompatible of a given element pair plotted on the x-axis. The peridotites show an extremely wide range in degree of depletion and form well-correlated depletion trends for upper mantle melting. Correlations are best for those element pairs having similar bulk D values (see the Sm-Nd, Sm-Eu and Dy-Er panels in fig. 2), and fall off slightly for elements with increasingly dissimilar bulk D's (see Zr-Ti and Sm-Yb). Some of the scatter in correlations may have been produced by melt impregnation. For instance, impregnation of low degree melts into highly depleted residues would cause points to fall off the depletion trends toward the more incompatible element, as shown in Figure 2. Melt impregnation trajectories for relationships among the HREE (e.g. Dy-Er) very closely overlap their well-correlated arrays (not shown for clarity), thereby indicating that melt impregnation has none to very little effect on the slopes or positions of these depletion trends.

The ln([A]) vs. ln([B]) depletion trends have been fit by a York (1996) two-error regression (fig. 2), assuming a blanket error of 10% for all points. The regressions have also been forced through the primitive upper mantle (PUM) coordinate of McDonough and Sun (1995); most regressions intersect the PUM coordinate without forcing, but this treatment becomes appropriate for some regressions with poor correlations (for example, see the Ce-Nd panel in fig. 2) and we find it necessary for obtaining robust output from the regression scheme.

The upper mantle is certainly not a unique composition (isotopic or otherwise), but rather a range of compositions reflecting variable amounts of depletion and/or reenrichment. A given peridotite's bulk trace element composition represents the culmination of melt depletion and enrichment over the entire history of that peridotite. Also, integrated within the trace element trends (fig. 2) is the net transfer of material out of the mantle by oceanic and continental crust generation as well as crustal recycling, in so far as the recycled material is well-mixed within the upper mantle. In this sense, some of the more complicated processes affecting upper mantle composition are incorporated into the abyssal peridotite trends, but are not required to explain the trace element data.

Knowing that much of the depletion observed in abyssal peridotites is caused by the most recent melting event, we need to define where today's *average*, *unmelted* upper mantle lies on the depletion trends. For this we need another constraint, the most obvious being the isotopic evolution of DMM: calculating the present-day Sm/Nd ratio from ¹⁴³Nd/¹⁴⁴Nd signatures in MORBs defines a unique position on the abyssal peridotite depletion trends.

2.2 Isotope Constraints

Although there has been much written on the homogeneity of MORBs relative to OIBs (Zindler and Hart, 1986; Hart, 1988; Hofmann, 1997), it is clear from the global database (fig. 3) that the MORB mantle is not one sole isotopic composition and by inference not one sole trace element composition. The variation in MORB 87Sr/86Sr, 143Nd/144Nd, and ²⁰⁶Pb/²⁰⁴Pb compared to all oceanic basalts is 18%, 35% and 47%, respectively. To address this issue of compositional heterogeneity in the upper mantle we derive three different trace element compositions for the MORB source by calculating present-day parent/daughter ratios from isotopic signatures and relating those ratios to the abyssal peridotite depletion trends. A study by Su and Langmuir (2003), who have filtered global MORB data for proximity to subduction zones and known plumes/hot spots (fig. 3; Table 1), provides an average MORB isotopic composition (Sr-Nd-Pb) as well as a standard deviation of the isotope data. We use the Su and Langmuir (2003) data analysis to define the following: 1) the average DMM composition, 2) an E-DMM based on isotopes that are 2σ enriched over the average, and 3) a D-DMM based on isotopes that are 2σ depleted from the average. The ¹⁷⁶Hf/¹⁷⁷Hf value of DMM is selected by averaging MORB values from Nowell et al. (1998), Chauvel and Blichert-Toft (2001) and Andres et al. (2002); 176Hf/177Hf values for E-DMM and D-DMM are based on correlations with 143Nd/144Nd (e.g. Chauvel and Blichert-Toft, 2001).

To derive present-day parent/daughter ratios, a depletion model must be assumed. It is common to use a two-stage evolution model with a single, instantaneous depletion event at a time (t) equal to the average age of the continental crust. An example from the Rb-Sr system, where T is the age of the Earth and parent/daughter ratios are for present-day is the following:

$$\left(\frac{^{87}Sr}{^{86}Sr}\right)_{DMM} = \left(\frac{^{87}Sr}{^{86}Sr}\right)_{0} + \left(\frac{^{87}Rb}{^{86}Sr}\right)_{RSF} \left(e^{\lambda T} - e^{\lambda t}\right) + \left(\frac{^{87}Rb}{^{86}Sr}\right)_{DMM} \left(e^{\lambda t} - 1\right) \tag{6}$$

However, because mantle depletion is largely the result of continental crust extraction, it is important to consider continental growth through time and the fact that it is not a single depletion event as modeled in the above equation. Although there is debate about continuous growth (Hurley and Rand, 1969) or constant volume with crustal recycling (Armstrong, 1968; Armstrong, 1981; Fyfe, 1978), somewhere in between (such as episodic growth) is probably the reality (McColloch and Bennett, 1994; Taylor and McLennan, 1995; Condie,

2000). Models suggesting continuous growth generally agree that the real increase in continental mass was at about 3 Ga (Taylor and McLennan, 1995). Models in favor of constant crustal volume suggest rapid (or at least effective) recycling of crust greater than 3 billion years old (e.g. Armstrong, 1981; Bowring and Housh, 1995). For our purpose, it is really preservation that is of issue; even though there are isotopic arguments for a 4.0 Ga depleted mantle of similar size and degree of depletion as the modern depleted mantle (e.g. Bowring and Housh, 1995; Vervoort et al., 1996), it is more important here that the abundance of that old crust is low compared to crust younger than 3.0 Ga. Isotopically, the recycling of old crust would make depletion appear to be younger on average and smear out (or even negate) the signal of ancient depletion.

Here we have improved on the classical methods by using a continuous transport, melt depletion model starting at 3 Ga. The equation for the Sr isotopic composition of DMM at time, t, as derived by Allègre (1969) and Hart and Brooks (1970) is:

$$\left(\frac{^{87}Sr}{^{86}Sr}\right)_{t} = \left(\frac{^{87}Sr}{^{86}Sr}\right)_{o} + \frac{\lambda\left(^{87}Rb/^{86}Sr\right)_{o}[1 - e^{-(\lambda + k)t}]}{\lambda + k} \tag{7}$$

where λ is the decay constant for ⁸⁷Rb and k is the difference in transport coefficients between Rb and Sr and essentially relates to the difference between bulk partitioning of the two elements. The initial conditions are values for bulk silicate Earth at the time when depletion begins (3.0 Ga). The parameter, k, is solved for by knowing the isotopic composition of bulk Earth at 3 Ga and DMM at time zero (i.e. present-day), as shown in Table 1. The parent/daughter variation through time is then:

$${\binom{87}{Rb}}^{86}Sr_{t} = {\binom{87}{Rb}}^{86}Sr_{t} e^{-(\lambda+k)t}$$
(8)

Figure 4 shows how this model, as used for the Rb/Sr system, gives a factor of ~2 lower estimate for the present-day Rb/Sr ratio of DMM, compared to the usual two-stage model. Decreasing the age of depletion requires lower Rb/Sr ratios for the same present-day ⁸⁷Sr/⁸⁶Sr ratio. In the same way that the Sr isotopes are used to constrain the Rb/Sr ratio of DMM, similar constraints result in determinations for Sm/Nd, Lu/Hf, and U/Pb (Table 1). The choice of depletion model does not greatly affect the Sm-Nd system, but is important for the whole suite of parent/daughter ratios.

Most important to the abyssal peridotite inversion is the present-day Sm/Nd ratio for DMM, since this identifies the present-day position of DMM on the mantle depletion arrays. Figure 5 shows how the Sm/Nd ratio of 0.411 in average DMM has a unique intersection with the Sm-Nd abyssal peridotite regression line, defining unique Sm and Nd concentrations of 0.239 ppm and 0.581 ppm, respectively. Given the absolute concentrations of Sm and Nd, we can derive the absolute concentrations of all the other trace elements that have been analyzed in abyssal peridotites (REE, Sr, Zr, Ti, Y; Table 2). Ultimately, estimates for every element concentration are obtained based on relationships with every other element. Standard deviations of various estimates for a given element are on the order of 1-2% for REE's and Ti and 5-6% for Sr, Zr and Ce, indicating that the depletion trends are internally very consistent; note that these standard deviations do not testify to the accuracy of the model. With other parent/daughter ratios, other pieces of the trace element pattern for DMM are estimated: Rb derives directly from the Sr; Hf derives from extrapolation to Lu from the REE's; Pb derives from the Ce/Pb canonical ratio (see below) and U follows (Table 2). The same procedure is applied with the E-DMM and D-DMM isotopic compositions.

It is possible to derive Th concentrations from Th/U ratios inferred from ²⁰⁸Pb/²⁰⁴Pb and ²⁰⁶Pb/²⁰⁴Pb isotopic evolution. However, it has been widely recognized that inferred Th/U ratios are generally much higher than observed Th/U ratios in MORBs; this has been termed the 'kappa conundrum' (Galer and O'Nions, 1985; Elliot et al., 1999; Tatsumoto, 1996). Since Th and U are both highly incompatible during mantle melting, with Th being slightly more incompatible, the Th/U ratios of MORBs should provide an upper limit to the source Th/U. Due to the extremely long half-life of ²³²Th (14 Byr), we interpret the kappa conundrum to be a problem where the Th-Pb system has not "caught-up" with the U-Pb system. For a more accurate representation of present-day upper mantle, we use Th/U ratios measured by careful U-Series studies on MORBs (Lundstrom et al., 1999; Sims et al., 2002; Sims et al., 2003) over the range in isotopic composition used to define our D-DMM through to E-DMM (Table 1).

2.3 Canonical Ratios and Volatile Contents

There are a few trace element ratios in MORBs (as well as many OIBs) that remain constant over variable degree of melting and variable isotopic composition. These ratios (Ce/Pb, Nb/Ta, Nb/U, and Ba/Rb) are termed "canonical" and their constancy is interpreted to be due to bulk partition coefficients being very nearly the same for the element pairs

during upper mantle melting, preventing significant fractionation (Hofmann and White, 1983; Hofmann et al., 1986; Jochum et al., 1997; Sims and DePaolo, 1997; Pfänder et al., 2002; Weyer et al., 2002). Hence, ratios in the melts are presumed to be identical to ratios of these elements in the source.

We employ these canonical ratios in order to complete the trace element pattern of DMM. It is clear though, from Figure 6, that some canonical ratios are better behaved than others; Nb/Ta, Nb/U and Ba/Rb all hover about their published values of 15.5, 47 and 11.3 respectively (Hofmann and White, 1983; Hofmann et al., 1986; Jochum et al., 1997; Pfänder et al., 2002; Weyer et al., 2002), whereas Ce/Pb ratios display much more scatter with a significant distribution of data higher than either published canonical values of 25 and 20 (Hofmann et al., 1986; Sims and DePaolo, 1997). As a mid-value of the MORB data, we use a Ce/Pb ratio of 30; if, in studies to come, it is determined that Ce/Pb should be lower, we recommend changing only the Pb concentration in DMM, as the U-Nb-Ta-La-Ce segment of the trace element patterns derived is consistent with MORB trace element characteristics (see section 3.2). This would imply that the derived U/Pb ratios from Pb isotopic inversions are not accurate – i.e. that U and Pb cycling in the upper mantle is more complicated than is assumed to be here. All other canonical ratios used are as previously published.

There is also a suite of canonical ratios involving the volatile elements H, C and Cl that are useful for estimating the budget of these elements in the upper mantle. Volatile elements play an important role in melt generation and transport, so the following estimates are crucial input parameters for modeling such processes. The compatibility of water has been shown to be similar to that of Ce, and MORBs have on average an H_2O/Ce ratio of ~200 (Michael, 1995). Using Ce = 0.550 ppm (Table 2), H_2O is calculated to be 110 ppm in DMM. Using a lower limit of 150 and upper limit of 250 for H_2O/Ce (Michael, 1995) along with our lower and upper estimates for Ce (Table 2), respectively, the range of water content in DMM is 70-160 ppm.

In a recent study on volatile undersaturated basaltic melt inclusions from the Siqueiros transform fault, Saal et al. (2002) have established two new working canonical ratios: CO- $_2$ /Nb at 239 \pm 46 and Cl/K at 0.0075 \pm 0.0025. Using our Nb min/max estimates from Table 2 and our K estimate from Table 3 (see section 3.1 below), average DMM has a CO $_2$ content of 36 \pm 12 ppm and a Cl content of 0.38 \pm 0.25 ppm. These estimates are about 2 times lower than those reported by Saal et al. (2002) due to their use of different trace

element abundances in DMM, but within error of recent estimates by Salters and Stracke (2004).

2.4 Final product

Combining the mantle depletion trends from the abyssal peridotites, parent/daughter ratios from the isotopic evolution of DMM, and a handful of canonical ratios, trace element patterns ("spidergrams") of average DMM, E-DMM (enriched from the average) and D-DMM (depleted from the average) have been derived (fig. 7a; Table 1 for isotope values; Table 2 for trace element compositions). Overall, the spidergrams are very smooth except for a dip in Sr along with an expected Pb anomaly (see Hofmann, 1988). The greatest difference between the three compositions is in the highly incompatible elements; for example, Rb and Ba vary by a factor of ~5, but Sm varies by only 30%.

Because it is difficult to assign errors to the estimated compositions, we have provided minimum and maximum estimates for trace element concentrations in the average DMM by using starting ages for the continuous depletion model at 2.5 and 3.5 Ga, respectively (see the grey lines on Figure 7b; Table 2). By changing the initial assumption of depletion age, we are effectively saying DMM is more or less depleted and all elements together move up or down the depletion arrays in proportion to their compatibilities during melting. Each of these patterns is internally consistent, and therefore not a true range for each element but rather a coherent solution for the assumptions made. It is not advisable to "mix and match" values from different patterns, since the elements behave as a system, not independently, and should be regarded as such. However, the Ba/Th ratios for the 2.5 and 3.5 Ga patterns are, respectively, too high and too low to generate realistic MORB compositions (see section 3.2) and support the idea that 3.0 Ga is an appropriate assumption for initiation of mantle depletion. As an aside, using a classical 1.8 Ga instantaneous depletion model results in a spidergram (not shown) very similar to that of the 3.5 Ga pattern but with Rb and Ba close to values for the 3.0 Ga (preferred) pattern.

The Salters and Stracke (2004) spidergram (fig. 7b) for average DMM, based on depleted MORB elemental ratios and isotopic evolution, is generally less depleted than the average DMM of this study. In particular, they show significantly higher estimates for the whole left side of the spidergram (Rb to Ce), as well as higher Zr, Hf and Y, than predicted even for our maximum estimate; this is most likely due to their use of an instantaneous (two-stage) depletion model for isotopic evolution instead of the gradual depletion model described in this paper.

Please note that in the following sections, only average DMM is discussed.

3. Physical and Chemical Properties of DMM

3.1 Modal abundances & major elements

Abyssal peridotites from all over the world show depletion trends not only in trace element content, but also in major element content in the form of modal abundances and mineral compositions (Dick et al., 1984; Dick and Natland, 1996; Hellebrand et al., 2002; Michael and Bonatti, 1985; Niu et al., 1997; Baker and Beckett, 1999). In general, these observations are supported by experimental and theoretical studies of peridotite melting (Kinzler and Grove, 1992a; Walter et al., 1995; Gaetani and Grove, 1998; Hirschmann et al., 1998). With increasing degree of melt depletion, olivine increases in modal proportion while orthopyroxene, clinopyroxene and spinel decrease. Al, Ca, Na, K and Ti are more incompatible during melting than Mg, Cr and Ni, so have the highest mineral and whole rock concentrations in the least depleted peridotites (e.g. Baker and Beckett, 1999).

Since we have established the systematics of incompatible trace elements in abyssal peridotites (see above), we use their relationship to modes to estimate the proportion of minerals in today's average DMM, then challenge the results with considerations of melt extraction and a comparison to the primitive mantle. Using the same data set as employed earlier (Suppl. Data Table 1), we have plotted bulk trace element concentrations against modal abundances in individual abyssal peridotites (see examples in fig. 8); note that these are not completely independent parameters since modes in part determine bulk trace element contents. By extrapolating the trends like those in Figure 8 to the DMM trace element concentrations (Table 2), we find DMM to be composed of 57% ol, 28% opx, 13% cpx and 2% sp (Table 3); orthopyroxene does not correlate with any trace elements and has been solved for by summation to 100%. These modes are very close to those found by extrapolating the modal abundance correlations for the South-West Indian Ridge abyssal peridotites (Dick, 1989) for which, without spinel, yields 60% ol, 30% opx and 10% cpx.

Constructing the bulk depleted mantle composition from mineral modes requires fairly precise knowledge of oxide abundances in the mantle minerals. Baker and Beckett (1999) have reduced mineral data for abyssal peridotites and provided algorithms for estimating major element compositions of minerals according to the modal abundance of olivine. Using the Baker and Beckett (1999) mineral compositions at 57% olivine and combining

the minerals in the given proportions, results in a bulk DMM having an Al₂O₃ content nearly that of the primitive upper mantle (PUM; McDonough and Sun, 1995), along with a very low Ca/Al ratio compared to PUM. Since Al is depleted during melt extraction and Al is more enriched than Ca in mantle melts (see Table 3), it is reasonable to expect the depleted mantle to have lower Ca and Al concentrations and a equal or higher Ca/Al ratio than that of PUM. This seems to be a problem only with Al abundance and is interpreted to be due to extrapolating mineral compositions beyond the limit of data used by Baker and Beckett (1999). For this reason, we find it more appropriate to use mineral compositions at 60% olivine, the lowermost (i.e. fertile) limit of their regressed data. In doing so, we calculate the major element composition of DMM as reported in Table 3.

Compared to PUM, estimated DMM has, as expected, lower SiO2, TiO2, Al2O3, CaO, Na₂O, and K₂O with higher MgO and Cr₂O₃ (Table 3). The TiO₂ content of 0.13 wt% is very close to the 0.12 wt% calculated from the trace element derivation (Table 2). The CaO/Al₂O₃ ratio is equal to that of PUM (0.80), and would increase to 0.86 by lowering the spinel abundance by only 0.5%. As previously mentioned, much of the trace element depletion in the upper mantle can be attributed to the continental crust. The question now is: can the newly estimated major element depletion also be attributed to the continental crust? The continental crust budget (see mass-balance section below and bulk continental crust of Rudnick and Fountain, 1997) does largely account for the low TiO₂ and K₂O, the most incompatible of the major elements, and partially accounts for the Na₂O. However, owing to the small mass of continental crust, major elements with less contrast between the crust and mantle cannot be balanced solely by the continents. As a possible solution, Table 3 shows a rudimentary mass balance of the DMM major elements in which 3% of primary MORB is subtracted from PUM (note that this is not a melting model). The major elements calculated in this way show striking similarity to the estimated DMM composition, implying that there is a isolated reservoir of oceanic crust somewhere within in the silicate Earth, as also suggested by Hauri and Hart (1997). A second estimate for Na₂O based on mass balance with both the continental crust and a MORB reservoir is also provided in Table 3 and is most likely more accurate than the original sum of mineral compositions.

To convert mineral modes into practical information for those who study abyssal peridotites, we have calculated the trace element composition of the constituent minerals in unmelted, average DMM by using Eqns. (1) and (2) with mineral/melt partition coefficients compiled by Kelemen et al. (2003). This effectively distributes the DMM trace element

budget among the minerals according to their relative affinity for incompatible elements. The resulting mineral compositions are reported as Suppl. Data Table 2

As a cautionary note, the reported modes and mineral compositions will only apply to the very shallow mantle, owing to variable mineral compositions and phases with pressure and temperature, but the average bulk composition of unmelted, upper mantle should be similar at any depth in this reservoir.

3.2 MORB generation

Here we present the average DMM trace element composition, so should be able to produce an average parental N-MORB. To model MORB generation, we first calculate bulk partition coefficients (D) from Eqn. (5) applied to the abyssal peridotite depletion trends. Because the slopes of the depletion trends (fig. 2) only supply information about *ratios* of D values, we initially have to assume a D for one element and then calculate the remainder in relation to that one. We have chosen a bulk D_{Nd} of 0.031, according to a compilation from Kelemen et al. (2003) for upper mantle melting, since the bulk compatibility of Nd is almost identical in both the spinel and garnet facies.

The set of partition coefficients resulting from the inversion of all abyssal peridotite depletion trends is reported in Table 2. Figure 9 shows the calculated D values compared to both garnet and spinel facies bulk D's from Kelemen et al. (2003); clearly the slope of the calculated D values very closely matches that of spinel facies melting and is far off the slope of garnet facies melting. Note that choosing a different D_{Nd} will change the absolute position but not the slope of the D values shown in Figure 9 (meaning the ratios of D's will not change). The above observation does not exclude a contribution of trace elements from small degree melts in the garnet facies, especially for elements with low D values; this may be why Sr and Ce are a little lower than the spinel facies prediction. For elements with higher bulk D's in the garnet facies, the majority of depletion will happen in the spinel facies, thus is reflected in the depletion arrays. If depletion for the middle to heavy REE's was significant in garnet melting, the trace element pattern across the REE would be steeper, even if the last D values to act on the peridotites were those of spinel melting. This is a straightforward effect of fractional melting on residues.

As an average MORB trace element pattern to be fit by our modeling, we use the parental (fractionation-corrected) N-MORB reported by Su and Langmuir (2003). A point generally agreed upon, and hence applied here, is that MORBs are aggregated fractional or near-fractional melts over a range of pressures, as initially confirmed by Sobolev and

Shimizu (1993). The highly incompatible elements are enriched ~16-fold over DMM, thus requiring 6% aggregated fractional melting of DMM. Figure 10 shows the excellent fit to N-MORB using the D values derived here and reported in Table 2.

The 6% degree of melting to generate MORB is on the low side of previous estimates ranging from 6-20% (Langmuir et al., 1992; Kinzler and Grove, 1992b). A higher F could be obtained if a greater fractionation correction is applied to the already-corrected parental N-MORB of Su and Langmuir (2003) since F is mainly limited by the concentration of highly incompatible elements (i.e. Rb, Ba, U, Th, Nb, Ta) in the modeled MORB; however, this requires generally higher D's to maintain the proper slope for M- to HREE. To get to F = 10%, at least another 30% fractionation is necessary and this seems unlikely. This low value for F, as compared to previous models, may be due to the fact that older estimates are based on anhydrous melting. Water content in the upper mantle is increasingly recognized as a very important aspect of mantle viscosity and convection (Hirth and Kohlstedt, 1996) as well and mantle melting (Hirth and Kohlstedt, 1996; Asimow and Langmuir, 2003; Asimow et al., 2004; Parman and Grove, 2004). To test the validity of 6% mean F, we apply our estimated concentration of H₂O to the recent model by Asimow et al. (2004) for hydrous mantle melting (the pHMELTS model).

A pHMELTS model run, using 110 ppm water, the major element compositions in Table 3, a potential temperature of 1360°C, a 2-D passive triangle melting regime and near-fractional melting, results in 7.0% mean melting to produce MORB and a crustal thickness of 6.3 km (P. Asimow, pers. comm., 2004). This confirms that the mean degree of melting (as defined by Plank et al., 1995) for hydrous models is generally on the low side of previous estimates from anhydrous models.

For average DMM, the derived D values (Table 2) translate into 2-3% melt depletion from primitive mantle using either non-modal or modal fractional melting models. Because average DMM is only 2-3% melt depleted, but average MORB is generated by 6% melting, one or all of the following is implied: 1) the degree of melting was lower in the past, 2) the mantle is a cannibal, recycling some crust and lithosphere back into itself, 3) depleted mantle has exchanged with the less depleted mantle, or 4) MORB generation is not the process that depletes the mantle.

4. Crust-Mantle Mass Balance

Classic works based on continental crust and the depleted mantle being sole complementaries calculate that from 30% to 70% of the mantle is depleted according to isotope and parent/daughter constraints [Allègre et al., 1983, refs. therein]. In the following paragraphs, we provide three different scenarios for the balance of chemical reservoirs in the silicate earth, with each scenario progressively involving additional reservoirs. We believe that using the entire trace element suite to assess the extent to which the silicate earth does or does not mass-balance is a robust way to calculate relative proportions of reservoirs. The mass of the continental crust is fixed to be 0.6% the mass of the silicate Earth. For each scenario, the best estimates of reservoir masses along with uncertainty in the estimates are derived from a Monte Carlo simulation with 10^5 trials in which element concentrations for each reservoir are considered as normal random variables with means given as PUM normalized concentrations and 1σ equal to 10%; in each trial, the relative reservoir masses are determined by a linear least-squares fit to the input parameters. Uncertainties are quoted as 2σ .

Scenario 1 is our own mass-balance involving only DMM and the continental crust, meaning in what proportion do DMM and the continental crust have to be added in order to sum back to primitive mantle? Our calculations indicate that 33±3% the mass of the silicate Earth has to be DMM, with the remainder being primitive mantle, to best balance the bulk continental crust composition as estimated by Rudnick and Fountain (1997). For this particular 'reconstituted' mantle, all elements sum to within 15% of the primitive mantle except Rb and Pb which are in excess by 28% and 21%, respectively. This scenario is no doubt overly simplistic. For one, there is no real evidence for the existence of a primitive mantle reservoir (see review by van Keken et al., 2002). However, there is evidence for a lower mantle that is less depleted than DMM, but depleted nonetheless; this reservoir is implied from many OIB isotopic arrays and termed FOZO by Hart et al. (1992). If a depleted lower mantle were accounted for here, the DMM reservoir would be significantly smaller than 33% in order to allow for the moderate depletion of the lower mantle.

Scenario 2 is the mass-balance of DMM, continental crust, and recycled oceanic crust. The idea of a recycled MORB component in the mantle is corroborated by the observation that the major element composition of DMM cannot be balanced by continental crust alone (see Table 3). At 43±6% DMM and 2.0±0.6% parental N-MORB of Su and Langmuir (2003), this scenario provides a better fit for most elements than does the first scenario; all

elements here are fit to within 8% except Rb and Pb with 20% and 15% excesses and Nb with a 17% deficit. The abundance of MORB cannot be increased greatly without having elevated M- to HREE that cannot be compensated by adding more DMM. However, we have not considered subduction zone alteration of oceanic crust, which almost certainly affects this mass balance.

Scenario 3 is the mass-balance of DMM, continental crust, recycled oceanic crust and an OIB source component (from Workman et al., 2004). To show upper limits on the sizes of enriched mantle components, the mass balance calculation has been made assuming 0% primitive mantle. The resulting balance is 74±5% DMM, 4.7±1.0% MORB and 21±5% OIB source; all elements are fit to within 15% except Rb and Nb with 23% excess and 21% deficit, respectively.

In each scenario 2 and 3, the fraction of MORB in the mass-balance is less than the mass of oceanic crust generated throughout Earth history (6-7% of the mantle mass, assuming constant rates for 4.55 Ga). As such, it is suggestive of the mantle being partially cannibalistic with respect to subducted oceanic crust. However, these mass balances do not include a depleted lithosphere, and could therefore be misrepresentative if oceanic crust and lithosphere are shown to be absolutely mechanically and thermally coupled.

5. Conclusions

Through abyssal peridotite trends, the isotopic composition of N-MORB and a few canonical ratios, we have constructed a major and trace element reference model for average, unmelted, depleted MORB mantle (DMM). From this DMM, parental MORB is generated by about 6% aggregated fractional melting as constrained primarily from the enrichment factor for the highly incompatible elements in parental MORB. Melting occurs mainly in the spinel facies mantle, but is most likely initiated within the uppermost garnet facies, as suggested by pHMELTS model runs. The water content inferred by the derived trace element composition is between 70 and 160 ppm, and is an integral ingredient in modeling mantle melting (Asimow et al., 2004) and mantle viscosity (Hirth and Kohlstedt, 1996; 2004). We have provided trace element compositions for minerals in the uppermost mantle which can be better estimated in the future with increasingly accurate information on mineral/mineral partitioning of trace elements and compositional variations of minerals with temperature and pressure.

The radiogenic heat production of DMM (from U, Th, and K) is only 15% that of the bulk silicate Earth at 2.43 x 10⁻⁹ Watts/m³, using a density of 3.20 g/cm³ and radiogenic heat production values from Durrance (1986). It takes ~33% of the mantle mass to be composed of DMM, the remaining being primitive mantle, to balance the continental crust alone, and ~43% if including an oceanic crust component. If there is no primitive mantle, the maximum masses of the oceanic crust and OIB reservoirs are 5% and 21%, respectively, if these enriched reservoirs are balanced purely by DMM.

Overall, the composition constructed here is a robust estimate for average upper mantle and can be used as an input parameter for a variety of models concerning mantle processes.

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Appendix 3-1.

Calculations for other melting models show that the fractional melting model used here to interpret residues of melting is a fine approximation for any number of more complicated (and maybe more realistic) melting models. Figure A1 plots melt depletion trends for Sm and Nd in residues of melting, starting with PUM of McDonough and Sun [14], and using equations for modal fractional, non-modal fractional, dynamic and non-modal dynamic melting as derived or summarized in Zou [23]. Residual porosity is chosen to be 0.5% for dynamic melting. Mineral/melt partition coefficients have been borrowed from Kelemen et al. [22] and the empirical coefficients of melting are from Niu [79] as follows: olivine = -0.17, orthopyroxene = 0.65, clinopyroxene = 0.47, spinel = 0.05. Bulk D values for Sm and Nd are taken from Table 2 in main text.

By definition, the two modal melting models are linear in Figure A1, with dynamic melting resulting in slightly less depletion for a given degree of melting (both curves end at 25% melting). Non-modal melting results in much greater depletion for a given degree of melting than modal melting and in slightly curved trajectories of melt depletion, but not severely enough to make a linear approximation (i.e. modal melting) invalid. For the upper few decades of melt depletion, the melting models are all nearly identical (fig. A1). Even at the point where they are clearly distinguishable from each other, the models all are contained within the scatter of the abyssal peridotite data (not shown on fig. A1 for clarity, but see figs. 2 and 5 above). Using modal fractional melting as an approximation for any of the other melting models will primarily lead to different D values than may be truly "Bulk D's", but will not change the ultimate estimate of DMM. For example, if the real melting style beneath ridges is non-modal dynamic melting, a linear fit by modal fractional melting will lead to estimates of 'effective' partition coefficients, meaning the 'non-modal' part of melting will be incorporated into the D outputs. Because we solve for relative partition coefficients (and do not assume them a priori), it is almost inconsequential which model is used given the resolution of the peridotite dataset. However, the estimate for the degree of melt depletion in DMM could be over-estimated if assuming an initial absolute D that is too high.

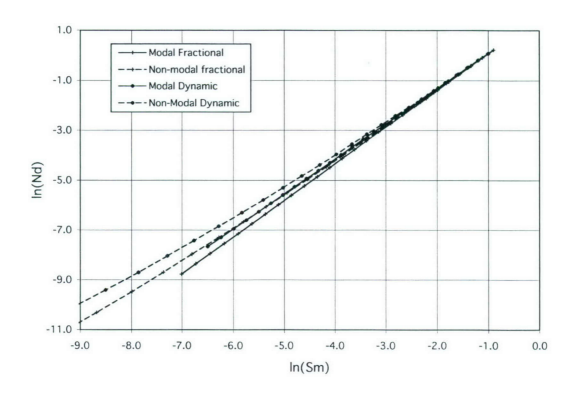


Figure A1.

Melting models showing depletion trends for Sm and Nd in residues of melt extraction. See text for details. Curves for modal melting models (fractional and dynamic) both end at 25% melt extraction. Non-modal fractional and dynamic curves are fall off the plot at 21% and 22% melting, respectively. Although not shown, the scatter in the abyssal peridotite data brackets all the melting model curves.

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Figures

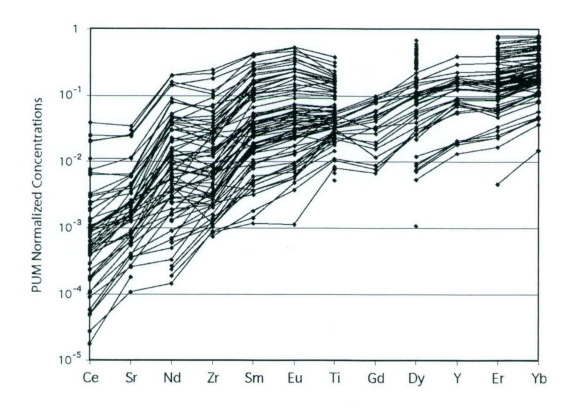


Figure 3-1. Calculated whole rock trace element compositions of abyssal peridotites. Data can be found in Suppl. Data Table 1; data sources as quoted in the text.

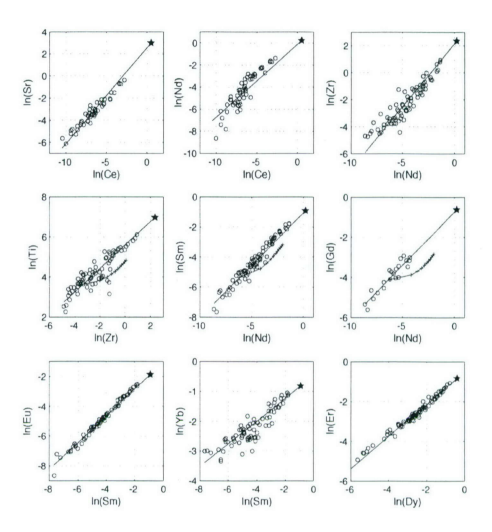


Figure 3-2. Co-variation diagrams for trace elements in abyssal peridotites, plotted as ln([A]) against ln([B]) in accordance with Eq. 4; the more incompatible element is on the x-axis for each plot. Star symbol is Primitive Upper Mantle (PUM; also Bulk Silicate Earth, BSE) of McDonough and Sun (1995). Solid line is the best-fit regression, forced through the PUM coordinate and with assigned errors of 10% for each point. The curved lines with crosses (in the middle three panels) are trajectories for impregnation of a 1% melt of DMM (see Fig. 7; D values from Kelemen et al., 2004) into a depleted mantle of composition indicated by the points of origin on the regression lines; ticks are in increments of 0.1% up to a total of 1% melt impregnation.

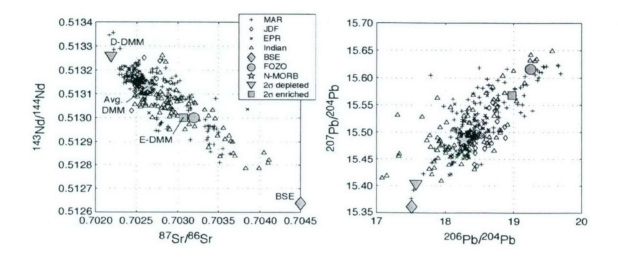


Figure 3-3. Two panels showing the Sr-Nd-Pb isotopic compositions of global MORB data obtained from the PETDB database (http://petdb.ldeo.columbia.edu/petdb/) MAR = Mid-Atlantic Ridge; JDF = Juan de Fuca Ridge; EPR = East Pacific Rise; Indian = Indian Ocean ridge systems; BSE = Bulk Silicate Earth; FOZO = Focus Zone (Hart et al., 1992); N-MORB and All-MORB are, respectively, the average composition for normal MORBs and unfiltered (all) MORBs according to Su and Langmuir (2003). The variation in N-MORB ⁸⁷Sr/⁸⁶Sr, ¹⁴³Nd/¹⁴⁴Nd, and ²⁰⁶Pb/²⁰⁴Pb compared to all of oceanic basalts is 18%, 35% and 47%, respectively.

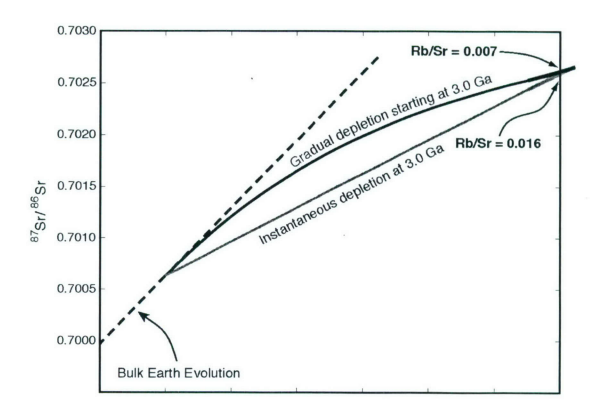


Figure 3-4. Plot showing examples of instantaneous depletion and continuous depletion models for the isotopic evolution of Sr in DMM. Note that the derived Rb/Sr ratio of today's upper mantle varies by a factor of about two between the models - the Rb/Sr ratio is defined by the slope of the evolution curves. Nd isotopic evolution by continuous depletion is not shown since it is essentially linear through time. See Table 1 for isotope evolution parameters and Eqs. 7 and 8 for continuous depletion model.

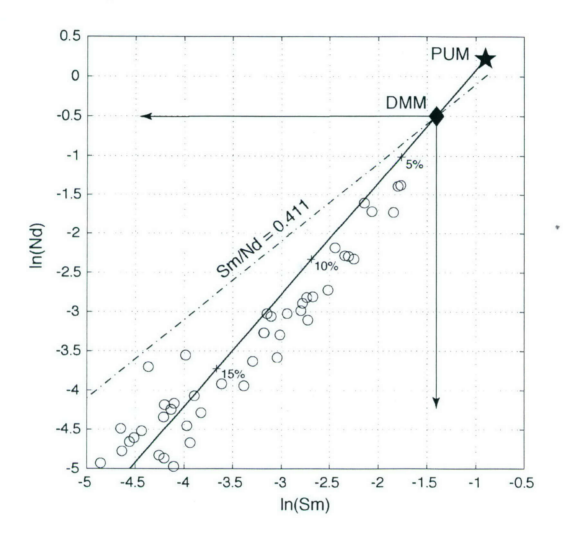


Figure 3-5. Plot of ln([Sm]) versus ln([Nd]) of abyssal peridotites showing the upper half of the mantle depletion array (see fig. 2 for full scale). Star symbol marks the PUM coordinate (i.e. BSE) of McDonough and Sun (1995). Crosses on the mantle depletion line mark percent melt extraction for fractional melting using bulk D values as reported in Table 2. Line of constant Sm/Nd of 0.411 is derived from the Nd isotopic evolution of DMM; the intersection of this line with the mantle depletion line marks today's average composition of DMM. From Sm and Nd concentrations, concentrations of all other elements reported for abyssal peridotites (see fig. 1) can be derived from the system of mantle depletion arrays like those shown in fig. 2.

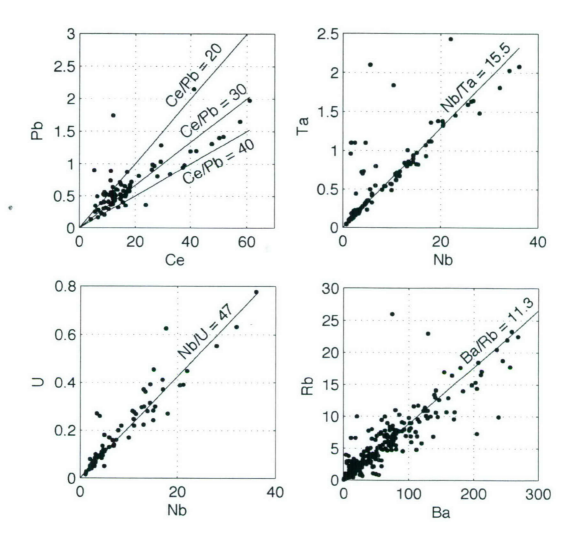


Figure 3-6. Four panels showing co-variation plots for elements in MORBs having canonical ratios. Data acquired from the PETDB database (http://petdb.ldeo.columbia.edu/petdb/) and reported as ppm. Literature values for canonical ratios (see text for references) are plotted for all panels except Ce-Pb, for which bounding and median values are shown.

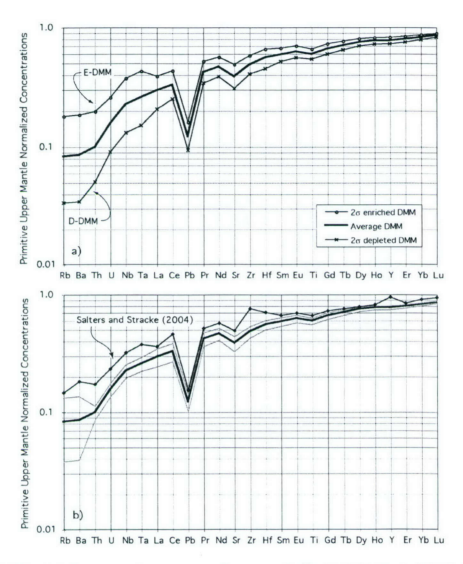


Figure 3-7. Panel a) shows trace element patterns for average Depleted MORB Mantle (DMM), E-DMM and D-DMM (E- and D- are, respectively, enriched and depleted over the average DMM, as based on a isotopic compositions reported in Table 1). Trace element compositions have been derived from abyssal peridotite depletion trends (see fig. 2), isotopic evolution based on the average N-MORB Sr-Nd-Pb-Hf isotopic compositions with a gradual depletion model starting at 3Ga (Table 1; Fig. 3), and canonical ratios (fig. 6). In panel b) the upper and lower grey lines result from using gradual depletion models starting at 3.5 Ga and 2.5 Ga, respectively, for the average DMM isotopic composition. The Salters and Stracke (2004) trace element pattern for average DMM is shown for comparison.

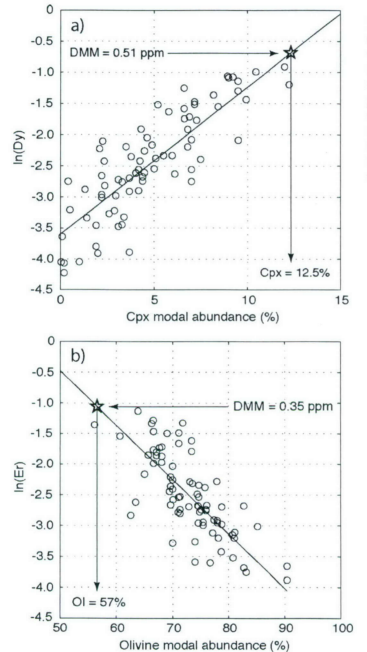


Figure 3-8. Modal abundances of a) clinopyroxene and b) olivine plotted with calculated (in the same way as for Figures 1 and 2) bulk trace element compositions of individual peridotites. Determination of the modal composition of DMM results from extrapolating the two-error regressions here to our best estimate for DMM's trace element concentrations (from Table 2).

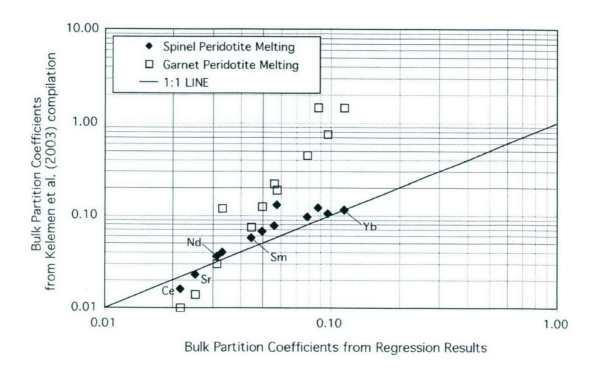


Figure 3-9. Bulk partition coefficients (D's) as obtained from abyssal peridotite trends (fig. 2, Table 2) plotted against both spinel and garnet facies bulk D values from Kelemen et al. (2003). Regression-based D values are calculated with Eqn. 5 and assuming $D_{Nd} = 0.031$. It is clear that the slope of the D values obtained reflect melting more in the spinel facies than the garnet facies, as the garnet facies is much steeper. Another choice for D_{Nd} will change the absolute concentrations, but not the slope, of the points plotted here.

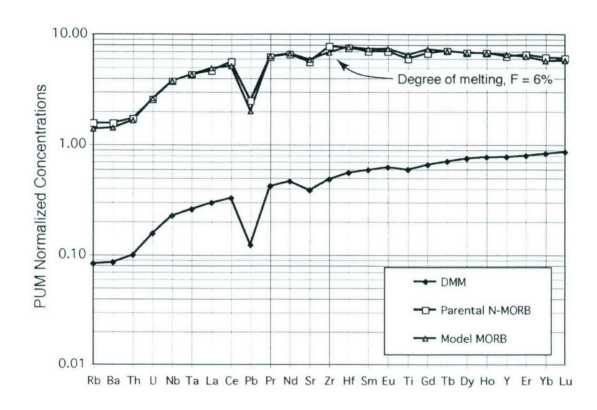


Figure 3-10. Plot showing the spectacular fit of parental N-MORB (from Su and Langmuir (2003) by 6% aggregated fractional melting of DMM (from this study) using Bulk D values in Table 2. The Rb, Ba, Th, U, and Pb for the parental MORB are taken from the N-MORB average of Hofmann (1988) and adjusted according to fractionation corrections provided by Su and Langmuir (2003).

Table 3-1. Isotopic constraints on the depleted MORB mantle (DMM).

	BSE ^a	DMM b	D-DMM	E-DMM	γ
$^{87}\mathrm{Sr/^{86}Sr}$	0.7045	0.70263	0.70219	0.70307	1.42×10^{-11}
143Nd/144Nd	0.512638	0.51313	0.51326	0.51300	6.54×10^{-12}
JH_{LL}/JH_{9L}	0.28276	0.28326	0.28350	0.28310	1.876×10^{-11}
$^{206}Pb/^{204}Pb$	17.511	18.275	17.573	18.977	1.55125×10^{-10}
$^{207}\text{Pb}/^{204}\text{Pb}$	15.361	15.486	15.404	15.568	9.8485 x 10 ⁻¹⁰
Rb/Sr°	0.0307	0.0065	0.0033	0.0111	
Sm/Nd	0.325	0.411	0.435	0.388	
Lu/Hf	0.239	0.369	0.439	0.325	
U/Pb	0.130	0.172	0.131	0.217	
Th/U d	3.9	2.5	2.2	3.0	

^a Age of Earth = 4.55 Ga.

^b Sr-Nd-Pb isotopic compositions taken from Su &Langmuir [28]; DMM is their average for MORBs far from plumes, D-DMM is 2σ depleted and E-DMM is 20 enriched over the average. See text for Hf isotope references.

^c Present-day parent/daughter ratios, calculated with a continuous depletion model starting 3 Ga. Decay constants (λ's) are summarized in Dickin [77].

^d Th/U ratios taken from U-Series disequilibrium studies on MORBs, as quoted in text.

Table 3-2. Trace element composition of DMM.

	Average				Enriched	Depleted			
	DMM		Ran	Range b	DMM	DMM		Calculated	
	(mdd)	Normalized a	Min	Max	(mdd)	(mdd)	Constraint	Bulk D°	10 (%)
Rb	0.050	0.083	0.023	0.079	0.108	0.020	Isotopes	0.00001	
Ba	0.563	0.085	0.256	968.0	1.219	0.227	Canonical Ratio	0.00012	
Th	0.0079	0.099	8900.0	0.0089	0.0157	0.0040	U-Series	0.001	
n	0.0032	0.156	0.0027	0.0036	0.0052	0.0018	Isotopes	0.0011	
NP	0.1485	0.226	0.1277	0.1671	0.2462	0.0864	Canonical Ratio	0.0034	
Ta	0.0096	0.259	0.0082	0.0108	0.0159	0.0056	Canonical Ratio	0.0034	
La	0.192	0.296	0.157	0.222	0.253	0.134	Extrapolated	0.01	
Ce	0.550	0.329	0.451	0.639	0.726	0.421	Regression	0.022	3.9
Pb	0.018	0.122	0.015	0.021	0.024	0.014	Canonical Ratio	0.014	
Pr	0.107	0.420	0.091	0.120	0.132	0.087	Interpolated	0.027	
PN	0.581	0.465	0.507	0.644	0.703	0.483	Regression	0.031	3.6
Sr	7.664	0.385	6.462	8.709	9.718	6.092	Regression	0.025	4.7
Zr	5.082	0.484	4.465	5.601	6.087	4.269	Regression	0.033	7.2
Hf	0.157	0.555	0.142	0.170	0.186	0.127	Isotopes	0.035	
Sm	0.239	0.588	0.217	0.256	0.273	0.210	Regression	0.045	2.2
Eu	960.0	0.624	0.088	0.102	0.108	980.0	Regression	0.050	2.5
Ti	716.3	0.594	666.5	756.1	792.0	650.0	Regression	0.058	2.9
PS	0.358	0.658	0.332	0.379	0.397	0.324	Regression	0.056	4.5
Tb	0.070	0.704	0.065	0.073	0.076	0.064	Interpolated	890.0	
Dy	0.505	0.749	0.480	0.525	0.543	0.471	Regression	0.079	7.3
Ho	0.115	0.772	0.110	0.119	0.123	0.108	Interpolated	0.084	
7	3.328	0.774	3.179	3.445	3.548	3.129	Regression	0.088	5.0
Er	0.348	0.795	0.334	0.359	0.369	0.329	Regression	0.097	8.7
Yb	0.365	0.827	0.353	0.374	0.382	0.348	Regression	0.115	8.1
Lu	0.058	0.859	0.056	0.059	0.060	0.056	Extrapolated	0.120	
		the straight in			0.,	-			

^a Normalized to primitive upper mantle (PUM) with values from McDonough and Sun (1995).

^b Minimum and maximum estimates for average DMM concentrations, based on assuming initiation of continuous depletion at 2.5 Ga and 3.5 Ga, respectively.

^c Bulk partition coefficients calculated from Eqn. 5, assuming D_{Nd} = 0.031 and solving for all other D's; 1 σ errors (%) are based on the variation of D estimates obtained from various regressions involving a given element. Values in italics are from Kelemen et al. (2003), except Rb, which is assumed, and U and Th, which are based on our own literature compilation. Underlined values are interpolated between regression outputs.

Table 3-3. Modal abundances and major element composition of DMM.

	Modal A	bundances	in DMM	(%):				
	Olivine	Opx	Cpx	Spinel				
	57	28	13	2				
	Mineral com	positions:	Срх	Spinel	Bulk DMM	PUM ^a	Primary N- MORB ^b	PUM minus 3% N- MORB
SiO ₂	40.70	53.36	50.61	Spiner	44.71	44.90	49.51	44.87
Al_2O_3	40.70	6.46	7.87	57.54	3.98	4.44	16.75	4.07
FeO*	10.16	6.27	2.94	12.56	8.18	8.03	8.05	8.05
MnO	0.14	0.12	0.09	0.16	0.13	0.13	0.14	0.13
MgO	48.59	30.55	16.19	19.27	38.73	37.71	9.74	38.68
CaO	0.05	2.18	19.52	17.27	3.17	3.54	12.50	3.27
Na ₂ O	0.03	2.10	17.52		0.13	5.54	12.50	3.21
1.420		0.05	0.89		$(0.28)^{c}$	0.36	2.18	0.30
Cr ₂ O ₃		0.76	1.20	10.23	0.57	0.38	0.07	0.39
TiO ₂		0.16	0.63	10.25	0.13	0.20	0.90	0.18
NiO	0.36	0.09	0.06	0.24	0.24	0.25	-	-
K ₂ O					0.006 d	0.029	0.065	0.028
P_2O_5					0.019 e	0.021	0.095	0.019
Total	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00
Mg # f	89.5	89.7	90.8	73.2	89.4	89.3	70.6	89.5
Cr # g				10.7				0,10
CaO/Al ₂ O ₃		0.34	2.48		0.80	0.80	0.75	0.80

^{*} Total Fe as FeO.

^a Primitive Upper Mantle (PUM) from McDonough and Sun (1995).

^b Primary N-MORB from averaged glass compositions in Presnall and Hoover (1987).

^c Value in parentheses is Na₂O estimated from the mass balance with continental crust (see text).

^d Calculated by inverting parental N-MORB at 0.1 wt% K₂O for 6% melting and assuming D_K = 0.0013.

^e Calculated by extracting 3% primary N-MORB (shown here) from PUM.

^f Mg # = molar ratio of Mg/(Mg+Fe²⁺); Mg # of N-MORB uses 90% total FeO as Fe²⁺.

^g C₅ # = molar ratio of C₇ (C₇+A₁).

 $^{^{}g}$ Cr # = molar ratio of Cr/(Cr+Al).

Supplementary Data Table 3-1. Abyssal peridotite modes and compositions

Sample	Location	Ridge Segment	Source Ref.			Modal	Data			Source Ref.
Number			(modes)	OL	OPX	CPX	SP	Plag	Sum	(Cpx 1E)
Vulc5:41-13(5)	Vulcan FZ	American-Antarctic	Dick (1989) G.S. Sp. Pub.	71.80	18.10	8.94	1.21	0.00	100.05	Johnson et al. (1990)
Vulc5:41-15(8)	Vulcan FZ	American-Antarctic	Sp.	63.90	22.70	12.00	1.44	0.00	100.04	Johnson et al. (1990)
Vulc5:41-30(3)	Vulcan FZ	American-Antarctic	Sp.	65.70	27.20	5.79	1.33	0.00	100.02	Johnson et al. (1990)
Vulc5:41-33(6)	Vulcan FZ	American-Antarctic	Sp.	68.10	21.00	9.93	96.0	0.04	100.03	Johnson et al. (1990)
Vulc5:41-45(7)	Vulcan FZ	American-Antarctic	Dick (1989) G.S. Sp. Pub.	68.10	22.30	8.45	1.14	0.04	100.03	Johnson et al. (1990)
Vulc5:34-56(3)	Bullard FZ	American-Antarctic	Sp.	74.80	17.10	7.50	09.0	0.00	100.00	Johnson et al. (1990)
Vulc5:35-1(4)	Bullard FZ	American-Antarctic		74.40	19.20	5.97	0.44	0.00	100.001	Johnson et al. (1990)
Vulc5:35-19(2)	Bullard FZ	American-Antarctic	Dick (1989) G.S. Sp. Pub.	70.00	22.10	68.9	1.09	0.00	100.08	Johnson et al. (1990)
Vulc5:35-22(4)	Bullard FZ	American-Antarctic	Dick (1989) G.S. Sp. Pub.	77.40	18.90	2.96	0.73	0.04	100.03	Johnson et al. (1990)
Vulc5:35-30(8)	Bullard FZ	American-Antarctic	Sp.	74.70	20.10	4.27	1.00	0.00	100.07	Johnson et al. (1990)
Vulc5:37-3(2)	Bullard FZ	American-Antarctic		82.70	13.90	2.35	0.05	1.05	100.05	Johnson et al. (1990)
A11107:40-6(5)	Bouvet FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	78.20	18.10	3.40	0.30	0.00	100.00	Johnson et al. (1990)
AII107:40-8(6)	Bouvet FZ	SW Indian		77.20	19.60	3.10	0.10	0.00	100.00	Johnson et al. (1990)
AII107:40-11(9)	Bouvet FZ	SW Indian		81.10	15.50	2.90	0.40	0.00	06.66	Johnson et al. (1990)
AII107:40-13(5)	Bouvet FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	78.70	18.80	2.00	0.50	0.00	100.00	Johnson et al. (1990)
AII107:40-27(5)	Bouvet FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	80.70	15.50	3.30	0.40	0.00	06.66	Johnson et al. (1990)
1011/76:56-57(4)	Islas Orcadas FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	09.69	24.50	4.32	1.66	0.00	100.08	Johnson et al. (1990)
1011/76:56-10(10)	Islas Orcadas FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	73.40	18.70	7.18	0.67	0.00	99.95	Johnson et al. (1990)
1011/76:56-54(4)	Islas Orcadas FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	00.69	21.70	8.20	1.14	0.00	100.04	Johnson et al. (1990)
1011/76:58-34(4)	Islas Orcadas FZ	SW Indian	Dick (1989) G.S. Sp. Pub.	70.00	25.90	3.51	0.62	0.00	100.03	Johnson et al. (1990)
PS86:6-37(3)	E. of Shaka FZ	SW Indian	Johnson et al. (1990)	99.99	21.85	10.43	1.13	0.00	26.66	Johnson et al. (1990)
Prot5:15-90(8)	Andrew Bain FZ	SW Indian	Johnson et al. (1990)	70.00	22.00	7.00	1.00	0.00	100.00	Johnson et al. (1990)
Prot5:19-2(8)	Pnc. Edward FZ	SW Indian	Johnson et al. (1990)	72.90	23.40	3.10	0.60	0.00	100.00	Johnson et al. (1990)
Prot5:29-26(6)	Discovery II FZ	SW Indian	Johnson et al. (1990)	63.50	28.60	7.00	06.0	0.00	100.00	Johnson et al. (1990)
Prot5:38-1(3)	Indomed FZ	SW Indian	Johnson et al. (1990)	69.40	24.80	5.10	0.70	0.00	100.00	Johnson et al. (1990)
RC27-9:6-3(7)	Atlantis II FZ	SW Indian	Johnson et al. (1990)	67.10	25.60	09.9	09.0	0.00	06.66	Johnson et al. (1990)
RC27-9:6-8(8)	Atlantis II FZ	SW Indian	Johnson et al. (1990)	09.99	25.80	09.9	0.80	0.00	08.66	Johnson et al. (1990)
RC27-9:25-142(6)	Atlantis II FZ	SW Indian	Johnson et al. (1990)	56.10	33.50	00.6	1.50	0.00	100.10	Johnson et al. (1990)
RC27-9:18-23	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	73.38	18.59	7.18	98.0	0.00	100.01	Johnson & Dick (1992)
RC27-9:18-45	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	77.92	19.39	2.25	0.43	0.00	66.66	Johnson & Dick (1992)
RC27-9:25-138	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	71.08	18.26	9.50	1.13	0.04	100.01	Johnson & Dick (1992)
RC27-9:30-32	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	42.50	42.10	15.09	0.46	0.00	100.15	Johnson & Dick (1992)
RC27-9:30-40	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	66.29	23.74	9.22	0.75	0.00	100.00	Johnson & Dick (1992)
RC27-9:34-58	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	70.94	25.37	3.31	0.37	0.00	66'66	Johnson & Dick (1992)
RC27-9:34-63	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	71.06	22.72	5.51	0.71	0.00	100.00	Johnson & Dick (1992)
RC27-9:35-49	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	75.96	19.53	4.05	0.45	0.00	66.66	Johnson & Dick (1992)
RC27-9:35-80	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	74.52	20.28	5.01	0.19	0.00	100.00	Johnson & Dick (1992)
RC27-9-44-1	Atlantis II FZ	SW Indian	Johnson & Dick (1992)	73.56	2133	4.64	0.47	000	100 00	Johnson & Dick (1992)

Supplementary Data Table 3-1, Part 1, Page 2

	Source Ref. (modes) Johnson & Dick (1992) Johnson & Dick (1990) Johnson & Dick (1990) Johnson & Dick (1990) Johnson & Dick (1990) Dick (1984) & unpubl.	OL 67.69 74.71 74.52 71.08 65.03	M OPX (Modal	Data			Source Ref. (Cpx TE)
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Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1993) 10-16 deg E SW Indian Johnson & Dick (1993) 10-16 deg E SW Indian Johnson & Dick (1990) 10-16 deg E SW Indian SW Indian Sw Indian Verna FZ Central Indian Sick (1984) & unpubl. Verna FZ Central Indian Verna FZ Central Indian Verna FZ Central Indian Sw Indian Johnson & Dick (1982) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II S SW Indian Johnson & Dick (1992) Atlantis II S SW Indian Johnson & Dick (1992) Atlantis II S SW Indian Johnson & Dick (1993) Atlantis II S SW Indian Johnson & Dick (1993) Atlantis II S SW Indian Johnson & Dick (1994) Argo FZ Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Argo F	Johnson & Dick (1992) Johnson & Dick (1990) Johnson & Dick (1990) Dick (1984) & unpubl.	74.52 71.08 65.03	22.24	2.09	96.0	0.00	100.00	Johnson & Dick (1992)
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Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1993) 10-16 deg E SW Indian Johnson & Dick (1994) Verna FZ Central Indian Johnson & Dick (1984) & unpubl. CIR axis 12° S SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1994) & unpubl. CIR axis 12° S Central Indian Johnson & Dick (1994) & unpubl. CIR axis 12° S Central Indian Johnson & Dick (1984) & unpubl. Argo FZ Central Indian Johnson & Dick (1984) & unpubl. Argo FZ Central Indian Johnson & Dick (1984) & unpubl. Argo FZ Central Indian Johnson & Unpubl. Argo FZ Central Indian Argo FZ Central Indian Johnson & Unpubl. Argo FZ Central Indian Johnson & Unpubl. Argo FZ Central Indian Argo FZ Central Indian Johnson & Unpubl. Argo FZ Central Indian Argo FZ	Johnson & Dick (1992) Johnson & Dick (1992) Johnson & Dick (1992) Johnson & Dick (1990) Johnson & Dick (1990) Johnson & Dick (1991) Dick (1984) & unpubl. Dick (1984) & unpubl. Dick (1984) & unpubl. Dick (1984) & unpubl.	65.03	18.26	9.50	1.13	0.04	100.01	Salters and Dick (2002)
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Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1993) Atlantis II FZ SW Indian 10-16 deg E SW Indian 10-16 deg E SW Indian 10-16 deg E Central Indian Verna FZ Central Indian Oick (1984) & unpubl. Oick (1984) & unpubl. Oick (1984) & unpubl. Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Johnson & Dick (1993) CIR axis 12° S Central Indian Johnson & Dick (1994) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian	Johnson & Dick (1992) Johnson & Dick (1990) Johnson & Dick (1992) Johnson & Dick (1990) Dick (1984) & unpubl.	90.89	23.78	7.29	0.79	0.07	66.66	Salters and Dick (2002)
Atlantis II FZ SW Indian Johnson & Dick (1990) Atlantis II FZ SW Indian Johnson & Dick (1991) 10-16 deg E SW Indian Johnson & Dick (1990) 10-16 deg E SW Indian Johnson & Dick (1990) 10-16 deg E SW Indian Verna FZ Central Indian Oick (1984) & unpubl. CIR axis 12° S Central Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II S Central Indian Johnson & Dick (1992) Atlantis II S Central Indian Johnson & Dick (1992) Atlantis II S Central Indian Johnson & Dick (1994) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Argo F	Johnson & Dick (1990) Johnson & Dick (1992) Johnson & Dick (1990) Dick (1984) & unpubl.	75.09	19.50	2.36	0.82	2.23	100.00	Salters and Dick (2002)
Atlantis II FZ SW Indian Johnson & Dick (1992) 10-16 deg E SW Indian Johnson & Dick (1999) 10-16 deg E SW Indian Verna FZ Central Indian Verna FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1994) Atrap FZ Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Johnson & Dick (1992) Johnson & Dick (1990) Dick (1984) & unpubl.	67.10	25.60	6.62	0.64	0.00	96.66	Salters and Dick (2002)
Atlantis II FZ SW Indian 10-16 deg E SW Indian Verna FZ Central Indian CIR axis 12° S Central Indian CIR axis 12° S SW Indian Atlantis II FZ SW Indian Johnson & Dick (1982) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ Central Indian Johnson & Dick (1992) Atlantis II FZ Central Indian Johnson & Dick (1992) Atlantis II FZ Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dic	Johnson & Dick (1990) Dick (1984) & unpubl.	75.96	19.53	4.05	0.45	0.00	66.66	Salters and Dick (2002)
10-16 deg E SW Indian Dick (1984) & unpubl. Verna FZ Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	09.99	25.80	19.9	08.0	0.00	18.66	Salters and Dick (2002)
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Vema FZ Central Indian Dick (1984) & unpubl. Central Indian CIR axis 12° S Central Indian Atlantis II FZ SW Indian Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian CIR axis 12° S Central Indian Argo FZ Central Indian CIR axis 12° S Central Indian CIR axis 12° S Central Indian CIR axis 12° S Central Indian Argo FZ Central Indian CIR axis 12° S Central Indian CIR axis 12° Centr	Dick (1984) & unpubl. Dick (1984) & unpubl. Dick (1984) & unpubl.	09.69	25.30	4.50	09.0	0.00	100.00	Hellebrand et al. (2002)
Verna FZ Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian CIR axis 12° S Central Indian Oick (1984) & unpubl. CIR axis 12° S W Indian Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Oick (1984) & unpubl. CIR axis 12° S Central Indian Oick (1984) & unpubl. CIR axis 12° S Central Indian Oick (1984) & unpubl. CIR axis 12° S Central Indian Oick (1984) & unpubl. Argo FZ Central Indian Oick (1984) & unpubl. Argo FZ Central Indian Oick (1984) & unpubl. Argo FZ Central Indian Argo FZ Central Indian Oick (1984) & unpubl.	Dick (1984) & unpubl.	09.69	25.30	4.50	09.0	0.00	100.00	Hellebrand et al. (2002)
Verna FZ Central Indian Verna FZ Central Indian Verna FZ Central Indian Verna FZ Central Indian CIR axis 12°S Central Indian Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12°S Central Indian Argo FZ Central Indian CIR axis 12°S Central Indian Argo FZ Central Indian Cir (1984) & unpubl. Central Indian Celeste FZ Central Indian Cir (1984) & unpubl. Central Indian Celeste FZ Central Indian Cele	Dick (1984) & unnubl	78.80	16.20	4.20	09.0	0.00	08.66	Hellebrand et al. (2002)
Verna FZ Central Indian Verna FZ Central Indian Verna FZ Central Indian CIR axis 12°S Central Indian CIR axis 12°S Central Indian CIR axis 12°S Central Indian Atlantis II FZ SW Indian Atlantis II FZ SW Indian Atlantis II FZ SW Indian SW Indian Atlantis II FZ SW Indian Oick (1984) & unpubl. SW Indian Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12°S Central Indian CIR axis 12°S Central Indian CIR axis 12°S Central Indian Argo FZ Central Indian Argo FZ Central Indian CIR axis 12°S Central Indian Argo FZ Central Indian CIR axis 12°S Central Indian Argo FZ Central Indian Argo FZ Central Indian CIR axis 12°S Central Indian CIR axis 12°S Central Indian Argo FZ Central Indian Cick (1984) & unpubl. Central Indian Contral Indian Cick (1984) & unpubl. Central Indian Contral Indi	industry of the state of the st	78.80	16.20	4.20	09.0	0.00	08.66	Hellebrand et al. (2002)
Vema FZ Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S W Indian Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	62.60	35.10	1.30	06.0	0.00	99.90	Hellebrand et al. (2002)
Verna FZ Central Indian CIR axis 12°S Central Indian CIR axis 12°S Central Indian CIR axis 12°S Central Indian Atlantis II FZ SW Indian Ohnson & Dick (1992) Atlantis II FZ SW Indian Atlantis II FZ SW Indian Ohnson & Dick (1992) CIR axis 12°S Central Indian CIR axis 12°S Central Indian Argo FZ Central Indian Argo FZ Central Indian Cick (1984) & unpubl. CIR axis 12°S Central Indian Argo FZ Central Indian Argo FZ Central Indian Cick (1984) & unpubl. Central Indian Argo FZ Central Indian Cick (1984) & unpubl. Central Indian Contral Indi	Dick (1984) & unpubl.	78.00	17.20	4.40	0.20	0.00	08.66	Hellebrand et al. (2002)
CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Dick (1984) & unpubl.	Dick (1984) & unpubl.	78.00	17.20	4.40	0.20	0.00	08.66	Hellebrand et al. (2002)
Atlantis II FZ SW Indian Dick (1984) & unpubl. Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	90.4	9.3	0.2	0.04	0	99.94	Hellebrand et al. (2002)
Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Maric Celeste FZ Central Indian Dick (1984) & unpubl. Maric Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.		9.3	0.2	0.04	0	99.94	Hellebrand et al. (2002)
Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Johnson & Dick (1992)	69.19	25.90	5.21	1.20	0.00	100.00	Johnson & Dick (1992)
Atlantis II FZ SW Indian Johnson & Dick (1992) Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Johnson & Dick (1992)		22.24	2.09	96.0	0.00	100.00	Johnson & Dick (1992)
Atlantis II FZ SW Indian Johnson & Dick (1992) CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl.	Johnson & Dick (1992)	74.52	20.28	5.01	0.19	0.00	100.00	Salters and Dick (2002)
CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Dick (1984) & unpubl.	Johnson & Dick (1992)	71.08	18.26	9.50	1.13	0.04	100.01	Salters and Dick (2002)
CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Maric Celeste FZ Central Indian Dick (1984) & unpubl. Maric Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	90.4	9.3	0.2	0.04	0	96.66	Hellebrand et al. (2002)
CIR axis 12° S Central Indian Dick (1984) & unpubl. CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Maric Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	75.50	21.70	2.20	0.70	00.0	100.10	Hellebrand et al. (2002)
CIR axis 12° S Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	75.50	21.70	2.20	0.70	0.00	100.10	Hellebrand et al. (2002)
Argo FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	83.10	16.70	0.10	0.10	0.00	100.00	Hellebrand et al. (2002)
Argo FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	70.10	23.40	6.10	0.40	0.00	100.00	
Argo FZ Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	71.30	20.90	7.00	0.80	0.00	100.00	Hellebrand et al. (2002)
Argo FZ Central Indian Dick (1984) & unpubl. Argo FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	71.30	20.90	7.00	0.80	0.00	100.00	Hellebrand et al. (2002)
Argo FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	74.60	20.80	3.70	0.90	0.00	100.00	Hellebrand et al. (2002)
Marie Celeste FZ Central Indian Dick (1984) & unpubl. Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	74.60	20.80	3.70	0.90	0.00	100.00	Hellebrand et al. (2002)
Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	76.70	19.50	3.70	0.10	0.00	100.00	Hellebrand et al. (2002)
	Dick (1984) & unpubl.	80.80	17.90	1.00	0.30	0.00	100.00	Hellebrand et al. (2002)
Marie Celeste FZ Central Indian Dick (1984) & unpubl.	Dick (1984) & unpubl.	82.70	17.10	0.04	0.14	0.00	86.66	Hellebrand et al. (2002)
ANTP 89 HD1 Marie Celeste FZ Central Indian Dick (1984) & unpubl. 71	Dick (1984) & unpubl.	71.30	27.80	0.40	0.40	0.00	06.66	-Hellebrand et al. (2002)

Supplementary Data Table 3-1, Part 1, Page 3

Sample	Location	Ridge Segment	Source Ref.			Modal	Data			Source Ref.
			(sanour)	OL	OPX	CPX	SP	Plag	Sum	(7 t vd)
ANTP 89 HD2	Marie Celeste FZ	Central Indian	Dick (1984) & unpubl.	76.10	21.20	2.20	0.30	0.00	08.66	Hellebrand et al. (2002)
ANTP 89 HD5	Marie Celeste FZ	Central Indian	Dick (1984) & unpubl.	73.10	20.50	4.90	0.50	1.00	100.00	Hellebrand et al. (2002)
ANTP 89 HD8	Marie Celeste FZ	Central Indian	Dick (1984) & unpubl.	75.80	19.90	3.80	0.30	0.00	08.66	Hellebrand et al. (2002)
ANTP 89 HD15	Marie Celeste FZ	Central Indian	Dick (1984) & unpubl.	85.10	14.40	0.50	0.04	0.00	100.04	Hellebrand et al. (2002)
ANTP 89 HD17	Marie Celeste FZ	Central Indian	Dick (1984) & unpubl.	81.00	16.90	1.90	0.10	0.00	06.66	Hellebrand et al. (2002)
ME33/2-31GTV	Green Rock Hill	Central Indian	Hellebrand et al. (2002)	70.00	28.00	1.40	09.0	NA	100.00	Hellebrand et al. (2002)
SO92-60GTV	Green Rock Hill	Central Indian	Hellebrand et al. (2002)	74.00	22.80	2.60	09.0	NA	100.00	Hellebrand et al. (2002)
S092-74GTV	Green Rock Hill	Central Indian	Hellebrand et al. (2002)	74.00	23.50	1.90	09.0	NA	100.00	Hellebrand et al. (2002)
G96-39/36	Romanche FZ	Mid-Atlantic	Tartorotti et al. (2002)	09.99	25.00	08.9	1.10	0.50	100.00	Tartorotti et al. (2002)
G96-39/36	Romanche FZ	Mid-Atlantic	Tartorotti et al. (2002)	09.99	25.00	08.9	1.10	0.50	100.00	Tartorotti et al. (2002)
895B 1R-1, 71-76	Site 895	East Pacific Rise	Dick and Natlland (1996)	87.70	11.30	0.31	0.62	0.12	100.05	Dick and Natlland (1996)
895C 1R-1, 28-32	Site 895	East Pacific Rise	Dick and Natlland (1996)	82.60	14.90	1.22	1.24	0.00	96.66	Dick and Natlland (1996)
895C 1R-1, 28-32	Site 895	East Pacific Rise	Dick and Natlland (1996)	82.60	14.90	1.22	1.24	0.00	96.66	Dick and Natlland (1996)
895C 3R-1, 35-39	Site 895	East Pacific Rise	Dick and Natlland (1996)	81.30	16.20	1.43	1.04	0.00	76.66	Dick and Natlland (1996)
895D 3R-1 116-120	Site 895	East Pacific Rise	Dick and Natlland (1996)	82.20	15.90	1.16	0.79	0.00	100.05	Dick and Natlland (1996)
895D 4R-3 97-101	Site 895	East Pacific Rise	Dick and Natlland (1996)	78.00	18.70	2.41	0.87	0.00	86.66	Dick and Natlland (1996)
895D 7R-2, 129-133	Site 895	East Pacific Rise	Dick and Natlland (1996)	82.20	15.60	1.71	0.48	0.00	66.66	Dick and Natlland (1996)

Supplementary Data Table 3-1, Part 2, Page 1

Sample	CPX Trace Element Content	Slement Con	tent													
Number	Concentration in ppm	ı in ppm														
	cbx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cbx	cpx	cpx	cpx	cpx
	Sc	^	Cr	La	Ti	Sr	Zr	Ce	PN	Sm	Eu	PS	Dy	E	4,4	>
Vulc5:41-13(5)	49.00	279.00	5637.00		2615.00	5.10	10.60	0.43	1.83	1.22	09.0		2.87	1.87	1.71	
Vulc5:41-15(8)	49.00	297.00	5925.00		2879.00	5.40	11.50	0.50	1.92	1.24	0.57		2.56	1.79	1.61	
Vulc5:41-30(3)	53.00	309.00	7530.00		1979.00	1.00	2.60	90.0	0.71	0.78	0.35		1.94	1.22	1.18	
Vulc5:41-33(6)	54.00	298.00	6759.00		1547.00	2.10	5.00	0.17	09.0	0.70	0.32		1.78	1.14	1.18	
Vulc5:41-45(7)	50.00	355.00	7928.00		1554.00	1.30	1.50	0.04	0.40	0.41	0.20		1.75	1.06	1.10	
Vulc5:34-56(3)	64.00	274.00	7579.00		196.00	09.0	09.0	0.01	80.0	0.15	80.0		0.87	0.49	0.32	
Vulc5:35-1(4)	70.00	300.00	7158.00		754.00	0.80	09.0	0.02	0.10	0.22	0.11		1.05	0.72	0.54	
Vulc5:35-19(2)	26.00	270.00	5876.00		1893.00	3.10	4.50	0.33	0.61	0.50	0.21		1.71	1.01	0.74	
Vulc5:35-22(4)	61.00	295.00	8150.00		886.00	1.80	0.30	0.01	0.04	0.12	0.07		0.82	0.64	0.57	
Vulc5:35-30(8)	54.00	321.00	8955.00		779.00	3.80	0.30	0.02	0.12	0.17	60.0		1.16	0.79	0.89	
Vulc5:37-3(2)	84.00	367.00	8275.00		1335.00	1.10	3.20	0.05	0.56	0.58	0.30		1.78	86.0	0.78	
AII107:40-6(5)	20.00	342.00	10227.00		244.00	1.20	0.10	0.01	0.02	0.05	0.02		0.55	0.46	0.57	
AII107:40-8(6)	70.00	285.00	6534.00		363.00	0.70	0.10	0.04	0.13	0.08	0.04		0.48	0.50	0.63	
AII107:40-11(9)	54.00	297.00	8392.00		443.00	0.50	0.10	0.01	0.02	0.05	0.03		0.70	0.57	0.70	
AII107:40-13(5)	53.00	291.00	8519.00		368.00	1.20	0.10	0.01	0.01	0.05	0.03		0.38	0.43	0.46	
AII107:40-27(5)	73.00	274.00	6518.00		326.00	0.80	0.20	0.02	0.04	0.07	0.03		0.52	0.52	0.53	
1011/76:56-57(4)	73.00	301.00	6042.00		1608.00	09.0	2.20	0.02	0.48	09.0	0.27		1.79	1.01	1.10	
1011/76:56-10(10)	44.00	278.00	6925.00		1320.00	09.0	1.40	0.02	0.47	0.48	0.30		2.12	1.57	1.57	
1011/76:56-54(4)	57.00	276.00	7793.00		1606.00	1.40	2.80	90.0	09.0	0.63	0.34		2.17	1.58	1.49	
1011/76:58-34(4)	52.00	278.00	6053.00		1113.00	20.20	1.10	1.55	09.0	0.34	0.18		1.46	1.01	69.0	
PS86:6-37(3)	52.00	293.00	6829.00		2674.00	5.40	11.00	0.87	2.16	1.37	0.56		2.65	1.70	1.67	
Prot5:15-90(8)	41.00	290.00	9462.00		1083.00	0.40	0.50	0.02	0.24	0.39	0.18		1.18	69.0	0.74	
Prot5:19-2(8)	78.00	258.00	7433.00		298.00	8.00	2.10	1.02	0.58	0.26	0.12		0.95	0.71	0.62	
Prot5:29-26(6)	74.00	276.00	7765.00		527.00	1.20	0.90	0.07	0.13	0.13	90.0		0.56	0.51	0.48	
Prot5:38-1(3)	62.00	282.00	7124.00		267.00	09.0	0.40	0.03	0.15	0.15	80.0		1.02	0.74	0.54	
RC27-9:6-3(7)	55.00	305.00	7051.00		1525.00	1.20	2.10	0.04	0.62	0.62	0.33		1.93	1.17	0.78	
RC27-9:6-8(8)	47.00	338.00	6959.00		1702.00	09.0	1.70	0.03	0.76	0.76	0.39		2.67	1.72	1.74	
RC27-9:25-142(6)	38.00	308.00	7334.00		1843.00	1.20	3.10	0.11	0.92	0.92	0.47		2.40	1.50	1.63	
RC27-9:18-23	26.00	326.50	5909.50		1875.50	1.01	2.50	90.0	0.75	0.80	0.39		2.22	1.33	1.39	
RC27-9:18-45	53.33	327.50	5617.50		1790.00	0.97	2.42	0.05	0.62	0.72	0.34		2.18	1.29	1.36	
RC27-9:25-138								0.12	0.98	0.88	0.40		2.17	1.29	1.35	
RC27-9:30-32			8673.33		1356.67	3.14	3.61	0.20	1.04	0.87	0.42		2.17	1.38	1.29	
RC27-9:30-40			7596.33		1747.00	1.29	3.06	0.10	0.97	06.0	0.43		2.62	1.70	1.59	
RC27-9:34-58			8690.00		537.00	0.44	0.20	0.02	0.04	0.13	90.0		98.0	0.61	0.64	
RC27-9:34-63			8771.50		00'.289	09.0	0.37	0.01	0.07	0.15	0.08		1.05	0.67	0.72	
RC27-9:35-49			8608.50		598.25	0.59	0.23	0.02	0.07	0.13	80.0		0.99	99.0	0.75	
RC27-9:35-80			79.7006		571.67	0.77	0.16	0.01	0.05	0.11	0.07		0.93	0.64	99.0	
RC27-9:44-1			8811.00		929.00	2.89	0.42	0.03	0.17	0.31	0.16		1.58	0.93	96.0	
RC27-9:46-1			7605.33		1521.33	0.71	1.29	0.03	0.43	0.67	0.30		2.35	1.47	1.54	
RC27-9:46-2			8395.50		1135.00	0.53	1.08	0.03	0.37	0.50	0.24		1.87	1.16	1.15	

Supplementary Data Table 3-1, Part 2, Page 2

Sample	CPX Trace Element Content	Element Co	ntent													
Number	Concentration in ppm	n in ppm														
	cbx	cpx	cbx	cbx	cbx	cbx	cpx	cbx	cbx	cbx	cbx	cbx	cpx	cpx	cpx	cpx
	Sc	Λ	Cr	La	Ti	Sr	Zr	Ce	PN	Sm	Eu	PD	Dy	Er	Yb	Υ
RC27-9 35-80					572.00	0.77	0.16	0.01	0.05	0.11	0.07		0.93	0.64	99.0	
RC 27-9 25-138	30.23		5507.37		2109.95	4.87	6.74	0.34	1.93	1.07	0.47		2.53	1.53	1.40	
RC27-9 30-44	29.04		7499.34		518.02	0.65	0.37	0.00	0.07	0.13	0.07		0.92	0.72	89.0	
RC27-9 34-62	22.87		4792.70		1103.61	1.06	1.69	90.0	0.56	0.49	0.19		1.53	1.02	0.95	
RC27-9 18-51	21.01		6182.36		96.099	0.58	0.43	0.01	0.17	0.25	0.11		1.05	08.0	69.0	
RC27-9 6-3					1535.00	1.19	2.32	0.04	0.47	0.57	0.33		1.64	1.23	1.18	
RC27-9 35-49					627.17	09.0	0.27	0.01	0.07	0.14	80.0		1.00	9.02	0.72	
RC27-9 6-8					1863.00	1.27	1.10	0.03	0.57	0.77	0.38		2.66	1.72	1.74	
PS86-6-26			5801.6	0.007	1325.7	1.740	2.483	0.093	0.834	0.611	0.267		1.842	1.125	0.994	
ANTP 134 HD3				0.0014	790.00	0.63	0.62	0.014	0.240	0.340	0.140	0.660	1.220	0.840	0.860	6.200
=				0.0023	00.899	1.35	1.00	0.016	0.250	0.250	0.120		0.860	0.620	0.560	00009
· ANTP 134 HD4				0.0020	657.00	0.88	0.54	0.036	0.260	0.270	0.130	0.680	1.090	0.730	0.830	5.700
Ε					628.00	1.07	1.00	0.035	0.310	0.300	0.120		0.780	0.550	0.570	5.800
ANTP 134 HD7					858.00	1.30	1.60	0.025	0.280	0.250	0.120		0.850	0.590	0.560	6.500
ANTP 134 HD8				0.0018	601.00	0.65	0.47	0.024	0.190	0.190	0.080	0.650	0.930	0.540	0.580	4.700
=					641.00	0.98	1.20	0.033	0.290	0.260	0.140		0.870	0.560	0.620	6.100
CIRCE 93 HD2				0.0140	965	1.19	0.40	0.028	0.060	0.120	0.065	0.420	0.750	0.640	0.720	5.400
=					825	1.60	92.0		0.120	0.140	0.070		0.640	0.510	0.490	4.800
CIRCE 93 HD4				0.0030	551.00	0.44	0.19	0.013	0.110	0.110	0.050	0.370	0.890	0.630	0.740	5.000
=					451.00	1.18	0.53	0.016	0.130	0.140	0.060		0.850	0.600	0.580	5.100
CIRCE 93 HD7				0.0150	601.00	3.24	3.50	0.095	0.500	0.460	0.210	0.840	0.840	0.440	0.530	4.200
ANTP 126 HD2				0.0000	391.00	0.54	0.21	0.049	0.040	0.070	0.040	0.250	0.730	0.610	0.660	4.500
ANTP 126-HD5				0.0100	388.00	0.47	0.17	0.045	0.060	0.070	0.040	0.340	0.740	0.620	0.750	4.600
=								0.041	0.000	0.000	0.050		0.610	0.470	0.570	
ANTP 126 D5				0.0050	516.00	0.58	0.17	0.020	0.050	0.130	090.0	0.390	0.940	0.710	0.840	5.200
=								0.020	0.080	0.120	0.080		0.760	0.540	0.620	
ANTP 84 HD11					130.00	0.19	0.07	0.002	0.005	0.009	0.003	0.060	0.290	0.290	0.400	2.000
ANTP 87 HD5				0.0025	259.00	0.25	80.0	0.002		0.030	0.020	0.130	0.420	0.460	0.490	2.800
ANTP 87 HD9				0.0050	345.00	0.48	0.11	0.007	0.020	0.040	0.040	0.220	0.560	0.470	0.590	3.800
ANTP 89 HD1				0.0024	913.00	0.72	0.78	0.013	0.210	0.300	0.130	0.760	1.330	0.840	0.950	6.800
ANTP 89 HD2				0.0019	767.00	0.40	0.28	900.0	0.060	0.160	0.100	0.550	1.220	0.840	0.910	6.700
ANTP 89 HD5				0.0010	791.00	0.27	0.21	0.005	0.130	0.230	0.130	0.730	1.380	0.950	1.080	7.500
ANTP 89 HD8				0.0030	817.00	0.29	0.26	9000	0.120	0.210	0.130	0.750	1.330	0.970	0.660	7.100
ANTP 89 HD15				0.0015	1419.00	0.63	0.95	800.0	0.170	0.210	0.130	0.710	1.260	0.930	0.920	7.200
ANTP 89 HD17				0.0140	575.00	0.33	0.13	0.013	0.012	0.060	0.040	0.240	0.640	0.570	0.720	4.200
ME33/2-31GTV				0.0020	523.00	0.39	91.0	0.004	0.040	0.060	0.030	0.230	0.610	0.430	0.580	3.400
SO92-60GTV				0.0050	500.00	0.42	0.13	0.017	0.030	0.090	0.040	0.270	0.600	0.430	0.580	3.500
SO92-74GTV				0.0013	460.00	0.29	0.12	0.003	0.020	0.040	0.020	0.180	0.390	0.330	0.330	3.200
G96-39/36	53.00	249.00	10400.00	0.01	353.00	0.24	0.41	0.05	0.14	0.11	90.0	0.32	1.03	1.02	1.00	8.30
G96-39/36	00.89	261.00	10210.00		445.00	0.44	08.0	0.07	0.22	0.19	0.07	0.47	1.37	1.28	1.07	10.80

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Sample	CPX Trace Element	_	Content													
Number	Concentration in ppm	mdd ui no														
	cpx	cpx	cpx	cbx	cbx	cpx	cpx	cbx	cbx	cpx	cpx	cpx	cbx	cpx	cpx	cpx
	Sc	>	Ċ	La	Ţ	Sr	Zr	Ce	PN	Sm	Eu	PS	Dy	Er	Yb	٨
895B 1R-1, 71-76		215.00	6387.00		147.00	0.500	0.400	0.013	0.111	0.167	0.036		0.135	0.161	0.212	0.800
895C 1R-1, 28-32		194.00	7511.00		150.00	0.400	0.200	0.010	0.023	0.021	900.0		0.126	0.165	0.229	1.000
895C 1R-1, 28-32		227.00	7165.00		173.00	0.700	0.500	0.004	0.015	0.010	0.002		0.018	0.033	0.071	0.900
895C 3R-1, 35-39		201.00	6740.00		184.00	0.800	0.300	0.010	0.018	0.014	0.007		0.132	0.168	0.204	1.000
895D 3R-1 116-120		201.00	6535.00		156.00	0.600	0.400	0.002	0.016	0.033	0.007		0.117	0.163	0.218	0.900
895D 4R-3 97-101		202.00	7599.00		208.00	0.400	0.300	0.015	0.033	0.046	0.009		0.142	0.186	0.231	0.800
895D 7R-2, 129-133		219.00	7137.00		163.00	0.900	0.600	0.019	0.044	0.063	0.020		0.137	0.193	0.215	0.900

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ordina or		Calculated of Hole Noen Links Element Collection										
Number	Concentration in ppm	mdd u										
	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock
	Ti	Sr	Zr	Ce	PN	Sm	Eu	PS	Dy	Er	Yb	Y
Vulc5:41-13(5)	333.80475	0.477810	1.889013	0.041236	0.180080	0.126382	0.064766	NaN	0.345861	0.266281	0.290742	NaN
Vulc5:41-15(8)	454.52213	0.676931	2.558331	0.064056	0.251890	0.170089	0.081226	NaN	0.401014	0.323966	0.333464	NaN
Vulc5:41-30(3)	193.39778	0.064311	0.454765	0.004055	0.050593	0.060998	0.029590	NaN	0.195579	0.156983	0.184595	NaN
Vulc5:41-33(6)	210.87544	0.218953	0.974458	0.018154	0.065806	0.080774	0.038518	NaN	0.238524	0.179634	0.217903	NaN
Vulc5:41-45(7)	190.63695	0.116698	0.277990	0.003698	0.038203	0.041610	0.021351	NaN	0.211703	0.154238	0.190196	NaN
Vulc5:34-56(3)	88.92315	0.047434	0.094848	0.000811	0.006681	0.013285	0.007417	NaN	0.091352	0.061979	0.049549	NaN
Vulc5:35-1(4)	72.72330	0.051398	0.089130	0.001331	0.006922	0.016433	0.008723	NaN	0.097137	0.083152	0.077864	NaN
Vulc5:35-19(2)	203.54483	NaN	0.762347	NaN	0.048688	0.042898	0.019115	NaN	0.180015	0.131182	0.116810	NaN
Vulc5:35-22(4)	60.64670	NaN	0.036028	NaN	0.001560	0.005336	0.003432	NaN	0.051074	0.054662	0.065567	NaN
Vulc5:35-30(8)	63.87800	NaN	0.041322	NaN	0.006320	0.009916	0.005683	NaN	0.089064	0.079375	0.115259	NaN
Vulc5:37-3(2)	79.98319	0.029496	0.306072	0.001423	0.017074	0.020325	0.011555	NaN	0.088741	0.068508	0.078051	NaN
AII107:40-6(5)	17.38195	NaN	0.011972	NaN	0.000860	0.002416	0.001054	NaN	0.036111	0.040605	0.067233	NaN
AII107:40-8(6)	24.56603	NaN	0.012012	NaN	0.005292	0.003706	0.002042	NaN	0.031006	0.044094	0.074267	NaN
AII107:40-11(9)	29.55918	NaN	0.010780	NaN	0.000735	0.002081	0.001363	NaN	0.040176	0.044713	0.075927	NaN
AII107:40-13(5)	21.50500	NaN	0.010929	NaN	0.000293	0.001741	0.001181	NaN	0.020011	0.032618	0.048700	NaN
AII107:40-27(5)	23.00745	NaN	0.022334	NaN	0.001630	0.003192	0.001483	NaN	0.031906	0.042827	0.059494	NaN
1011/76:56-57(4)	136.21770	0.029390	0.340106	0.001039	0.026537	0.037048	0.018227	NaN	0.148235	0.110254	0.151336	NaN
(01)/16:56-10(10)	143.68200	0.045738	0.224013	0.001569	0.038108	0.041487	0.027278	NaN	0.220162	0.198368	0.242752	NaN
1011/76:56-54(4)	193.18173	0.121979	0.507286	0.005385	0.055633	0.062116	0.035270	NaN	0.255496	0.224265	0.252542	NaN
1011/76:58-34(4)	81.68029	NaN	0.159788	NaN	0.028696	0.018563	0.010921	NaN	0.111857	0.105044	0.091659	NaN
PS86:6-37(3)	378.88240	0.591089	2.234232	NaN	0.248582	0.165690	0.070616	NaN	0.371060	0.278872	0.318693	NaN
Prot5:15-90(8)	117.23475	0.030080	0.084838	0.001557	0.019408	0.033862	0.016566	NaN	0.125355	0.090230	0.117445	NaN
Prot5:19-2(8)	41.47878	NaN	0.281158	NaN	0.024671	0.012701	0.006526	NaN	0.065913	0.067646	0.077174	NaN
Prot5:29-26(6)	56.86989	NaN	0.169556	NaN	0.010921	0.011856	0.005871	NaN	0.064176	0.073039	0.081874	NaN
Prot5:38-1(3)	50.43465	0.034112	0.063142	0.001794	0.009480	0.010459	0.006044	NaN	0.092789	0.086936	0.078769	NaN
RC27-9:6-3(7)	157.09406	0.086446	0.365686	0.003003	0.048714	0.052837	0.030101	NaN	0.206209	0.156391	0.125961	NaN
RC27-9:6-8(8)	176.41230	0.043251	0.298787	0.002255	0.059791	0.064866	0.035641	NaN	0.285895	0.230472	0.281370	NaN
RC27-9:25-142(6)	238.16168	0.117452	0.698379	0.011207	0.097846	0.105252	0.057386	NaN	0.337447	0.257944	0.322856	NaN
RC27-9:18-23	205.39304	0.076585	0.401810	0.004864	0.060371	0.069088	0.035423	NaN	0.229707	0.167119	0.213806	NaN
RC27-9:18-45	108.48071	0.026237	0.273391	0.001499	0.019916	0.026962	0.014197	NaN	0.121835	0.102058	0.148670	NaN
RC27-9:25-138	NaN	NaN	NaN	0.012668	0.101734	0.095896	0.045467	NaN	0.273349	0.190775	0.236186	NaN
RC27-9:30-32	250.10150	0.504841	1.097559	0.033140	0.178155	0.157957	0.079335	NaN	0.459141	0.342989	0.350348	NaN
RC27-9:30-40	224.96774	0.126492	0.596380	0.010499	0.101109	0.099481	0.050138	NaN	0.341781	0.264367	0.290273	NaN
RC27-9:34-58	37.91421	0.017329	0.027618	0.000645	0.001975	0.006618	0.003745	NaN	0.063615	0.061655	0.082820	NaN
RC27-9:34-63	63.84291	0.035978	0.057649	0.000821	0.004608	0.010866	0.006497	NaN	0.097016	0.078776	0.105135	NaN
RC27-9:35-49	46.66126	0.026737	0.030126	0.000712	0.003563	0.007249	0.004594	NaN	0.073162	0.064547	0.094525	NaN
RC27-9:35-80	49.37056	NaN	0.023042	NaN	0.003047	0.007242	0.004712	NaN	0.078473	0.068860	0.090057	NaN
RC27-9:44-1	77.80027	NaN	0.059662	NaN	0.009395	0.019480	0.010867	NaN	0.129194	0.099020	0.129307	NaN
RC27-9:46-1	139.50056	0.041328	0.213146	0.001656	0.027880	0.047583	0.023285	NaN	0.219081	0.177224	0.228200	NaN
RC27-0-46-2	101104	170010	200000									

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Number	Concentration in ppm	mdd u										
	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock
	Ti	Sr	Zr	Ce	PN	Sm	Eu	PS	Dy	Er	Yb	٨
RC27-9 35-80	49.39935	NaN	0.023037	NaN	0.003047	0.007242	0.004712	NaN	0.078473	0.068860	0.090057	NaN
RC 27-9 25-138	280.07410	0.483614	1.237919	0.034323	0.201335	0.116933	0.053253	NaN	0.319749	0.226115	0.246283	NaN
RC27-9 30-44	68.50270	NaN	0.074477	NaN	0.008000	0.014191	0.008134	NaN	0.124150	0.115096	0.127148	NaN
RC27-9 34-62	121.73116	0.083303	0.298858	0.005141	0.046956	0.044630	0.018671	NaN	0.170892	0.139780	0.156208	NaN
RC27-9 18-51	41.07591	0.016437	0.049331	0.000301	0.005740	0.009570	0.004594	NaN	0.059938	0.064351	0.075293	NaN
RC27-9 6-3	158.66144	0.085964	0.404637	0.002996	0.037181	0.049032	0.030350	NaN	0.175126	0.165196	0.189989	NaN
RC27-9 35-49	48.91665	0.027076	0.035147	0.000704	0.003379	0.007527	0.004564	NaN	0.073531	0.063412	0.091149	NaN
RC27-9 6-8	193.28625	0.091674	0.193443	0.002057	0.044900	0.065512	0.035070	NaN	0.285448	0.231091	0.282083	NaN
PS86-6-26	209.94257	0.224078	0.576202	0.012326	0.112972	0.086754	0.039571	NaN	0.301579	0.213940	0.214725	NaN
ANTP 134 HD3	65.44163	0.032111	0.094844	0.000756	0.013785	0.021785	0.009802	0.049248	0.104509	0.094553	0.121370	0.827773
	55.33545	0.068810	0.152974	0.000864	0.014359	0.016019	0.008402	NaN	0.073670	0.069789	0.079032	0.801071
ANTP 134 HD4	52.47788	0.040347	0.066668	0.001697	0.013025	0.014861	0.007671	0.042214	0.077545	0.067483	0.101165	0.658673
=	50.16150	0.049059	0.123459	0.001673	0.015530	0.016513	0.007081	NaN	0.055491	0.050843	0.069475	0.670229
ANTP 134 HD7	45.48473	0.027633	0.243477	0.000635	0.008436	0.009662	0.005607	NaN	0.056391	0.058782	0.071795	0.774008
ANTP 134 HD8	48.35045	0.031254	0.059301	0.001203	0.009987	0.010963	0.004951	0.042333	0.069205	0.052023	0.072930	0.560548
=	51.56845	0.047121	0.151406	0.001654	0.015243	0.015002	0.008664	NaN	0.064740	0.053950	0.077960	0.727520
CIRCE 93 HD2	38.15369	0.005052	0.024670	0.000150	0.000408	0.001268	0.000849	0.006294	0.017105	0.025849	0.051830	0.358780
:	32.61844	0.006792	0.046874	NaN	0.000817	0.001479	0.000914	NaN	0.014596	0.020598	0.035273	0.318915
CIRCE 93 HD4	33.65233	0.011938	0.022949	0.000386	0.003600	0.004262	0.002196	0.017730	0.051899	0.052460	0.083562	0.528592
	27.54483	0.032017	0.064016	0.000476	0.004255	0.005424	0.002635	NaN	0.049566	0.049962	0.065495	0.539164
CIRCE 93 HD7	23.31880	0.016102	0.289229	0.000660	0.004669	0.006660	0.003901	0.018271	0.026202	0.023476	0.044677	0.326327
ANTP 126 HD2	38.14694	ZaZ	0.033987	NaN	0.002901	0.005514	0.003372	0.022144	0.072359	0.076053	0.100951	0.655516
ANTP 126-HD5	41.73910	NaN	0.028118	NaN	0.004820	0.006027	0.003643	0.032273	0.077590	0.079808	0.117619	0.689754
	NaN	NaN	NaN	0.003175	0.007230	0.007749	0.004553	NaN	0.063959	0.060500	0.089391	NaN
ANTP 126 D5	39.29340	0.024315	0.022731	0.000888	0.002365	0.006904	0.003484	0.024149	969290.0	0.068310	0.105310	0.615510
	NaN	NaN	NaN	0.000888	0.003784	0.006373	0.004646	NaN	0.054733	0.051954	0.077729	NaN
ANTP 84 HD11	9.54850	NaN	0.008795	NaN	0.000233	0.000470	0.000171	0.003634	0.020420	0.027224	0.049354	0.233081
ANTP 87 HD5	12.46761	0.003562	0.007673	0.000029	NaN	0.000728	0.000572	0.004157	0.017481	0.029623	0.046490	0.246829
ANTP 87 HD9	13.23075	0.002142	0.009177	0.000045	0.000179	0.000566	0.000733	0.004746	0.017417	0.025150	0.049806	0.295489
ANTP 89 HD1	38.95086	0.007600	0.093206	0.000180	0.003704	0.007449	0.004073	0.027643	0.064111	0.064795	0.101489	0.671045
ANTP 89 HD2	45.72279	0.010807	0.032785	0.000177	0.001949	0.006145	0.004347	0.026066	0.070364	0.069150	0.101955	0.702862
ANTP 89 HD5	68.00623	0.014540	0.029816	0.000281	0.007688	0.014917	0.009067	0.053685	0.115175	0.101880	0.146693	0.967107
ANTP 89 HD8	61.28521	0.012387	0.033482	0.000270	0.005743	0.011236	0.007576	0.046483	0.095605	0.092671	0.123606	0.837610
ANTP 89 HD15	60.25429	0.005313	0.075515	0.000081	0.002081	0.003558	0.002670	0.016563	0.040635	0.049247	0.077290	0.559808
ANTP 89 HD17	32.25031	NaN	0.013079	NaN	0.000329	0.001954	0.001468	0.009576	0.031536	0.040660	0.073180	0.399382
ME33/2-31GTV	27.71900	0.008035	0.020937	96000000	0.001109	0.002096	0.001244	0.010707	0.035602	0.037538	0.067640	0.368867
SO92-60GTV	32.27500	NaN	0.016549	NaN	0.001118	0.003912	0.001955	0.014326	0.038200	0.038378	0.068827	0.389639
S092-74GTV	26.59375	0.007120	0.014708	0.000082	0.000612	0.001478	0.000850	0.008427	0.022471	0.027626	0.037387	0.338638
G96-39/36	37.58568	0.017735	0.072515	ZaZ	0.011242	0.009549	0.005560	0.031106	0.111241	0.137029	0.161878	1.281852
(196-39/36	XYIXY IV					NI-NI	11-11	000000	010000	01010	0000000	1 //1000

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Sample	Calculated Wi	Calculated Whole Rock Trace Element Content	ce Element C	ontent								
Number	Concentration in ppm	in ppm										
	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock	bulk rock
	Ti	Sr	Zr	Ce	PN	Sm	Eu	PS	Dy	Er	Yb	\
895B 1R-1, 71-76	6.25485	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.003564	0.007269	0.016187	0.056440
895C 1R-1, 28-32	8.01188	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.005033	0.010026	0.020876	0.084906
895C 1R-1, 28-32	9.24036	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.000719	0.002005	0.006473	0.076416
895C 3R-1, 35-39	10.07630	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.005766	0.010970	0.019496	0.089130
895D 3R-1 116-120	8.01060	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.004756	0.010129	0.020193	0.077586
895D 4R-3 97-101	13.23400	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.008033	0.014810	0.025271	0.082114
895D 7R-2, 129-133	9.05873	NaN	NaN	NaN	NaN	NaN	NaN	NaN	0.006265	0.012929	0.020940	0.081931

Supplementary Data Table 3-2. Mineral compositions in average DMM.

		Mine	ral Modes		
	ol	opx	cpx	sp	sum
	0.57	0.28	0.13	0.02	1
	Miner	al Trace elemen	t compositions	(ppm)	
	ol	opx	срх	sp	sum ^a
Rb	-	0.005	0.371	-	0.050
Ba	-	0.062	4.197	-	0.563
Th	-	0.0001	0.061	-	0.008
U	-	0.0002	0.024	-	0.003
K	-	0.541	389.6	-	50.800
Nb	0.058	0.167	0.443	0.575	0.149
Ta	0.004	0.011	0.029	0.037	0.010
La	0.0002	0.026	1.414	0.016	0.192
Ce	0.0005	0.137	3.930	0.027	0.550
Pb	0.00002	0.005	0.129	0.001	0.018
Pr	0.0002	0.034	0.744	0.003	0.107
Nd	0.002	0.194	4.044	0.013	0.581
Sr	0.004	1.312	56.11	-	7.664
Zr	0.657	6.574	20.28	11.50	5.082
Hf	0.013	0.131	0.836	0.228	0.157
Sm	0.004	0.109	1.585	0.003	0.239
Eu	0.002	0.056	0.611	0.001	0.096
Ti	101.8	1018	2715	1018	716.3
Gd	0.007	0.239	2.209	0.004	0.358
Tb	0.003	0.051	0.415	0.001	0.070
Dy	0.026	0.396	2.916	0.010	0.505
Но	0.010	0.096	0.633	0.003	0.115
Y	0.751	3.266	15.25	0.147	3.328
Er	0.036	0.363	1.735	0.012	0.348
Yb	0.086	0.376	1.615	0.017	0.365
Lu	0.016	0.065	0.235	0.003	0.058

Chapter 4:

Volatile and Trace Elements in Basaltic Glasses from Samoa: Implications for Water Distribution in the Mantle

Abstract

We report volatile (H₂O, CO₂, F, S, Cl) and trace element data for submarine alkalic basalt glasses from the three youngest Samoan volcanoes, Ta'u, Malumalu and Vailulu'u. Most samples are visibly sulfide saturated, so have likely lost some S during fractionation. Cl/K ratios (0.04 – 0.15) extend to higher values than pristine MORBs, but are suspected to be partly due to source differences since Cl/K roughly varies as a function of ⁸⁷Sr/⁸⁶Sr. There are no resolvable differences in the relative enrichment of F among sources, and compatibility of F during mantle melting is established to be nearly identical to Nd. Shallow degassing has affected CO₂ in all samples, and H₂O only in the most shallowly erupted samples from Vailulu'u. Absolute water contents are high for Samoa (0.63 – 1.50 wt%), but relative enrichment of water compared to trace element enrichment is low. H₂O/Ce (58 – 157) and H₂O/La (120 – 350) correlate inversely with ⁸⁷Sr/⁸⁶Sr compositions (0.7045 – 0.7089). This leads us to believe that, because of very fast diffusion of hydrogen in olivine, recycled lithospheric material with high initial water content will lose water to the drier ambient mantle during storage within the inner Earth. The net result is the counter-intuitive appearance of greater dehydration with greater mantle enrichment.

1. Introduction

Volatile elements such as hydrogen, carbon, fluorine, sulfur and chlorine track not only the melt phase in magmatic systems, but also the gas and fluid phases. Because of their ability to decouple from lithophile elements, especially at or near the Earth's surface, these elements provide a possible fingerprint of processes that are not actively recorded by the standard suite of incompatible trace elements (e.g. Sr, Rb, U, Th, and the rare earth elements). They have been used extensively to study volatile-driven arc magmatism, but have not been as widely applied to ocean island volcanism. It is now well established that the distribution of water in the mantle (primarily in nominally anhydrous minerals) strongly determines the locations and extents of mantle melting (Asimow and Langmuir, 2003) along with mantle rheology (Hirth and Kohlstedt, 1996; 2003), so it is important to understand the abundance of water in all mantle materials.

Lavas from the Samoan islands and seamounts have unique geochemical characteristics compared to other ocean island basalts (OIBs) as well as mid-ocean ridge basalts (MORBs), including extreme ⁸⁷Sr/⁸⁶Sr (up to 0.7089), low ¹⁴³Nd/¹⁴⁴Nd (down to 0.5125), and highly trace element enriched lavas (Wright and White, 1986; Farley et al., 1992; Workman et al., 2004). These characteristics define the Enriched Mantle 2 reservoir (EM2; Zindler and Hart, 1986). EM2 has classically been interpreted as oceanic crust and terrigeneous sediment that had been recycled back to the mantle through subduction zones (e.g. Weaver, 1991). Upon closer inspection, the standard recycling model of ocean crust/sediment fails as an explanation for producing EM2, due to smooth trace element patterns, low 187Os/188Os ratios, incorrect Pb isotope compositions, and high 3He/4He (>8R-A) in Samoan lavas (Workman et al., 2004). Instead, the origin of EM2 has been modeled with the ancient formation of metasomatised oceanic lithosphere, followed by storage in the deep mantle and return to the surface in the Samoan plume. Although Samoa is strongly EM2 in character, other mantle reservoirs, such as HIMU (high ²³⁸U/²⁰⁴Pb mantle; Zindler and Hart, 1986) and PHEM (Primitive Helium Mantle; Farley et al., 1992), are thought to contribute to the Samoan lavas, and hence in one location there is extreme compositional heterogeneity.

Here we report a study of volatile element concentrations in Samoan lavas in order to assess their relative enrichment compared to other trace element and isotope signatures, and to use them as potential clues to the origin of compositional variations in the mantle. We

find that water enrichment (identified by H₂O/Ce and H₂O/La ratios) in EM2 lavas has an inverse relationship to lithophile element enrichment (e.g. high Rb/Sr and ⁸⁷Sr/⁸⁶Sr ratios). The model for the formation of EM2 (Workman et al., 2004) would not result in fractionation of water from the lithophile elements. However, the fast diffusion of water through mantle minerals (Mackwell and Kohlstedt, 1990) provides a very effective means of losing water from EM2 to a water-poor ambient mantle during the long-term (~2.5 Ga) storage of EM2 – to the point of there being very similar bulk water contents (<100 ppm) in EM2 and the Depleted MORB Mantle (DMM). This finding leads credence to the idea that enriched mantle plumes are indeed "hot-spots" (Morgan, 1971; Sleep, 1990; Campbell and Griffiths, 1990) and not "wet-spots" (Schilling, et al., 1980; Green and Falloon, 1998).

2. Background

2.1. Geological and Geochemical Setting

The Samoan islands and seamounts are centered on 14°S latitude, stretch from 169-177.5°W longitude (fig. 1), and rest on the outer rim of the geophysical "superplume" of the South Pacific (Su et al., 1994). Seismic tomography images from beneath Samoa show a slow velocity anomaly extending into the lowermost mantle (Montelli et al., 2003), supporting a deep origin for upwelling mantle material feeding the Samoa hotspot. The tectonic setting of Samoa is complicated by having the northern termination of the Tonga Trench only 100 km to the south of the island chain. Hart et al. (2004a) have given a synthesis of the regional tectonic history and how plume-trench interaction has influenced the distribution of Samoan volcanism, specifically the *en-echelon* topographic ridges seen in Figure 1. Hart et al. (2004a) also established, from K-Ar dating and isotope geochemistry of seamounts extending to the west of Savai'i, that the Samoan plume has been active for at least 25 million years, producing a 1700 km long volcanic chain.

The easternmost, leading-edge Samoan volcano is the seamount, Vailulu'u (fig. 1); it rises from a 5000 m deep seafloor to a summit depth of 590 m and rests 45 km east of its nearest neighbor, Ta'u Island (Hart et al., 2000). Vailulu'u has been shown to be both volcanically and hydrothermally very active (Hart et al., 2000; Hart et al., 2003; Staudigel et al., 2004; Konter et al., 2004). The power output from Vailulu'u's summit crater is estimated to be 610-760 MW, the equivalent of 20-100 Mid-Ocean Ridge black smokers;

Mn export is estimated at 240-300 kg/day, or the equivalent of about 10 black smokers (Staudigel et al., 2004; Hart et al., 2003). The presence of ³He anomalies in the water column show that at least this volatile element is actively degassing from upwelling magmas beneath Vailulu'u (Staudigel et al., 2004). We suspect that the hydrothermal circulation of seawater and degassing of magmas observed at Vailulu'u is representative of all the Samoan volcanoes at this stage of growth.

One goal of this study is to understand volatile enrichments/depletions as a function of mantle source composition. Each of these Samoan volcanoes has a distinct and well-characterized geochemical signature, discussed in detail by Workman et al. (2004). In summary, Malumalu displays the most extreme EM2 signature yet documented from ocean island basalts, with ⁸⁷Sr/⁸⁶Sr up to 0.7089 and extreme trace element enrichment. Ta'u shows some of the least enriched lavas from Samoa, with Sr-Nd-Pb isotopic values that are close to the mantle component PHEM (Primitive Helium Mantle of Farley et al. [1992]; ⁸⁷Sr/⁸⁶Sr = 0.7045); for Ta'u this corresponds to ³He/⁴He up to 18 R_A and a lesser degree of trace element enrichment. Vailulu'u lavas are of intermediate enrichment and are thought to contain a small component of HIMU mantle, giving Vailulu'u the highest ²⁰⁶Pb/²⁰⁴Pb (19.4) in Samoa.

2.2. Samples and Volcano ages

Glass samples from the three youngest Samoan volcanoes, Vailulu'u, Malumalu, and Ta'u, have been taken from glassy rims of pillow basalts dredged during the 1999 AVON2/3 cruise aboard the R/V Melville. In this paper, we report data on samples obtained from 11 dredges ranging from 780 to 4170 meters deep – 6 from Vailulu'u, 2 from Ta'u and 3 from Malumalu (see Table 1). All Ta'u and Malumalu dredges are from a similar depth of 2300 – 3600 meters, whereas the shallowest and deepest dredges are from Vailulu'u.

All samples have some vesicles, most with less than 10% by volume and none over about 30%, indicating that these samples have experienced volatile exsolution during eruption. All dredges except 72 (Vailulu'u) and 75 (Ta'u) have sulfides present in nearly every sample (see Table 1); these occur as 10-50 µm spheres and are visible under reflected light in the glass chips that were analyzed. Micro-phenocryts of olivine, spinel, and clinopyroxene are common within the glasses. In a few of the olivine phenocrysts, melt inclusions have been fortuitously exposed, so these were analyzed for their volatile contents

to compare with volatile abundances in the matrix glass, but they have not been properly studied for major elements and mineral-melt equilibration.

U-series data have been collected for samples from each Vailulu'u and Malumalu dredge (Sims and Hart, 2004), including the following samples studied here: 63-13, 70-01, 71-02, 71-09, 72-02, 73-01, 73-03, 73-12, 76-01, 78-01. All show ²³⁰Th/²³⁸U excesses and are interpreted to be less than 300 thousand years old. The ²³⁰Th/²³⁸U excesses shown from Malumalu are similar to those on Vailulu'u, suggesting that Malumalu is not significantly older than Vailulu'u. Two samples from Vailulu'u (63-13 and 70-09) show ²¹⁰Pb disequilibria are interpreted to be less than 100 years old (Hart et al., 2000).

The oldest K-Ar age from Tau Island is 0.3 Ma (McDougall, 1985). The youngest volcanic series on Tau (Faleasao) is probably younger than 37,000 years, based on ¹⁴C ages of coral inclusions in these volcanics (Hart, unpublished). Additionally, there was an underwater eruption just west of Tau in 1866 (see description in Keating, 1992).

3. Analytical Techniques

Multiple glass chips from the quenched rims of dredged pillow basalts were hand-picked and mounted with epoxy for ion microprobe analyses. Volatile abundances (H_2O , CO_2 , F, S and Cl) were determined by triplicate analyses on the Cameca IMS 6f ion microprobe at the Department of Terrestrial Magmatism with a 5-10 nA Cs⁺ primary beam. See Hauri et al. (2002) for a full description of the mircobeam method. Trace elements were determined on the same area of glass using the same instrument, but with an O^- primary beam. Analytical uncertainties are $\pm 10\%$ for H_2O and $\pm 5\%$ for all other elements. All calibrations were made against mafic glass standards.

For Sr isotope analyses, glass chips were hand-picked and leached for 1 hour in 6N HCl. Sr was separated by a standard cation exchange procedure described in Taras and Hart (1987). Isotopic analyses were carried out on the NEPTUNE multi-collector ICP/MS at W.H.O.I.; ⁸⁷Sr/⁸⁶Sr values are corrected to 0.71024 for NBS 987 and carry a 2σ precision equal to ±25 ppm (Hart et al., 2004b).

4. Results

Volatile and other trace element concentrations for Samoan glasses are listed in Table 1. Major element abundances of the glasses were analyzed by electron microprobe at MIT and

have been previously published (Workman et al., 2004). Sr, Nd and Pb isotope measurements have been made on a few whole rock samples (for which we now report glass data) and are also published in Workman et al. (2004). New ⁸⁷Sr/⁸⁶Sr measurements on glass samples are reported in Table 1, and range from 0.70452 to 0.70841, within the previously established range for whole rock analyses of Samoan lavas (Workman et al., 2004). The ⁸⁷Sr/⁸⁶Sr compositions of glass and whole rock pairs for individual samples have been found to be within analytical error (Jackson et al., unpubl.), so using ⁸⁷Sr/⁸⁶Sr values from "mixed material" is inconsequential to our data analysis.

4.1. Trace elements and fractionation correction

All the glasses presented here are evolved beyond clinopyroxene saturation, as identified by: 1) low Mg#'s (molar percent of Mg/[Mg+Fe⁺²]), ranging from 39 to 61; 2) positive correlations between Mg# and CaO; 3) presence of clinopyroxene phenocrysts in whole rock samples; and 4) liquid lines of descent predicted by the pMELTS model (see fig. 2). They are all alkali basalts, with SiO₂ from 45.5 to 48.9 wt% and total alkalis (K₂O+Na₂O) from 3.2 to 6.1 wt%. The glasses are generally further along on fractionation trends than their whole rocks (fig. 2) due to the exclusion of phenocrysts in the glasses. Mg#'s of aphyric whole rocks are equal to or slightly higher than their glasses, but some glasses come from picritic samples with Mg#'s up to 81-84.

Trace element patterns of the glasses closely parallel whole rock trace element patterns (from Workman et al., 2004) of each volcano. La/Sm and Sm/Yb for glass and whole rock pairs are usually within analytical error, showing that the fractionation of olivine and clinopyroxene has had little effect on incompatible element ratios. Hence, using the data in this way (as elemental ratios) would be the most reliable in assessing primary melt characteristics.

La/Sm values of the glasses range from 3.3 to 7.5 and correlate positively with ⁸⁷Sr/⁸⁶Sr, a proxy for source enrichment (i.e. Ta'u has the lowest and Malumalu has the highest La/Sm; not shown). Sm/Yb, on the other hand, does not correlate with enrichment, and the three volcanoes overlap in values from 2.9 – 5.6. For comparison, N-MORB La/Sm and Sm/Yb ratios are each ~1.0 (Hofmann, 1988). The steepness of Samoan rare-earth element slopes is a strong indication of melting within the stability field of garnet lherzolite (Hauri et al., 1994; Salters et al., 2002), as to be expected with melting beneath an old tectonic plate where the bottom of the lithosphere (e.g. Li et al., 2004) is near the garnet to spinel transition (Kogiso et al., 1998; Robinson and Wood, 1998; Klemme and O'Neill, 2000).

In order to estimate the compositions of primitive Samoan magmas, we have corrected the raw glass data for crystal fractionation by using the pMELTS trends (Ghiorso et al., 2002) to calculate the remaining liquid mass as a function of Mg#. The degree of fractionation ranges from 17% to 66%, with the majority of lavas falling within the range of 30 - 46% fractionated. Figure 2 shows that Malumalu and Ta'u lavas lie along very similar fractionation lines. Vailulu'u lavas are on a separate trend at higher CaO, with shallow dredges (63, 68, 70 and 73) being more fractionated on average than deep dredges (70 and 71).

The same fractionation corrections have been applied to the volatile contents, assuming that crystal fractionation happened at vapor under-saturated conditions. Figure 3 shows the fractionation-corrected abundances of volatile elements and a select group of trace elements, listed in the order of increasing compatibility during mantle melting. Concentrations are normalized to primitive upper mantle (PUM); most values for PUM are taken from McDonough and Sun (1995) except for CO₂ and H₂O, which are calculated assuming that PUM has a CO₂/Nb ratio of 239 (Saal et al., 2002) and an H₂O/Ce ratio of 200 (the value for FOZO observed by Dixon et al., 2002). This plot allows for a brief overview of the relative enrichments of volatile elements compared to the trace elements, suspending (for the moment) any interpretation about loss of volatiles by degassing. For all volcanoes, the negative anomalies for CO2, H2O, and S indicate that these volatiles are less enriched in the lavas than lithophile elements of similar compatibility; the opposite is true for the positive anomaly at Cl for most of the Vailulu'u lavas. F is the only volatile element that is consistently "conformable" with the lithophile elements in all samples. In the following sections, we attempt to discern which of these signals are primary and how they can be used to understand the composition and evolution of the mantle.

4.2. Water and carbon dioxide in glasses

The Samoan lavas show a large range in raw H_2O (0.63 – 1.50 wt%) and CO_2 (6 – 233 ppm) contents (fig. 4a; Table 1). Such water contents are highly elevated over MORBs (fig. 4b), but are very similar to Hawaii's submarine lavas from Loihi Seamount (Dixon and Clague, 2001) and the North Arch lava field (Dixon et al., 1997), and to melt inclusion data from the Austral Islands (Lassiter et al., 2002). Water in OIB lavas is generally lower than in arc lavas (up to \sim 6 wt%) but in the same range as back-arc basin lavas (e.g. Newman et al., 2000, for the Marianas system).

The presence of vesicles (1-30 vol%; Table 1) in every sample indicates that these lavas were all vapor saturated during eruption onto the seafloor. Some vapor loss can be accounted for by correcting for vesicle gases (i.e. the manifestation of closed system degassing). As a test for the severity of water loss, vesicle abundances were determined in 3 of the most vesicle-rich samples, 73-01, 73-03, and 78-03, by point-counting standard-sized thin sections. By volume, these contain 31%, 20% and 24% vesicles, respectively. Adding equilibrium vapor compositions back into the glasses, following the procedure described in Dixon et al. (2002), results in increases in bulk water content on the order of 0.03 to 0.05 wt%; yet these samples still have the lowest water abundances. Because this level of correction is within error of the analyses, we chose not to make vesicle corrections for all samples in the Samoan suite. It is likely that the volume of vesicles present in the glasses at eruption does not represent the total amount of degassing experienced by these lavas; in other words, we think it is likely that these lavas have lost gas via open-system degassing with bubble loss, prior to their final eruption and emplacement on the seafloor.

 $\rm H_2O\text{-}CO_2$ solubility models (Stolper and Holloway, 1988; Dixon et al., 1995; Dixon, 1997) show that $\rm CO_2$ is much more insoluble than $\rm H_2O$ in basaltic magmas. As a result, with Samoan water contents, vapor compositions during open-system degassing are almost entirely $\rm CO_2$ until pressures drop below ~100 bars (ocean depth of 1000 meters) (Newman and Lowenstern, 2002). In Figure 4a, the arrow at the end of the degassing trend marks the sharp change from $\rm CO_2$ -dominated to $\rm H_2O$ -dominated vapor compositions; above this point, open-system degassing results in nearly negligible water loss.

In Figure 4c, calculated equilibrium saturation pressures (using the Dixon [1997] model for H₂O-CO₂ saturation in alkali basalts) are plotted with collection depth. Lavas that lie very close to the 1:1 line erupted close to their dredge depth. Lavas below the 1:1 line are interpreted to have erupted more shallowly and flowed down slope; lavas above the line experienced incomplete degassing (i.e. rapid quenching). Note that some scatter is because dredges are along flanks of the volcanoes and hence may traverse 100's of meter of depth contours (see fig.1); the quoted dredge depth is the average for each dredge and may not represent every sample accurately.

As indicated by carbon dioxide concentrations and pressures of equilibration, both the most and the least degassed samples are from Vailulu'u: the deep dredges (71 and 72 from 4200 m and 3800 m deep, respectively) show water contents up to 1.50 wt%, and the shallowest dredges (63, 68, 70, and 73 at or near 1000 m) have water contents down to 0.63 wt%. The low water in the shallow dredges is the result of degassing a water-rich vapor (fig.

4a). Ta'u and Malumalu lavas all have high enough pressures of equilibration (fig. 4c) to preclude loss of significant water by open system degassing.

In summary, degassing has affected CO₂ in all samples, and H₂O only in the most shallowly-erupted samples from Vailulu'u. Additional support for minimal water loss is that fact that the most melt inclusions have water contents that are identical (within error) to their host matrix glasses, but with equal or higher CO₂ (Tables 1 and 2; the one exception is a melt inclusion from sample 71-03, which has lower CO₂ and H₂O than its matrix glass). One melt inclusion from sample 72-12 shows the highest pressure of vapor saturation at 818 bars, indicating a minimum crystal fractionation depth of 1.6 km within the oceanic crust (3840 meters dredge depth plus 1600 meters igneous crust at a rock density of 2800 kg/m³).

4.3. Water and carbon dioxide in primary magmas

After fractionation-correction (see section 4.1), we estimate primary magmas to have H_2O in the range of 0.4 - 1.1 wt%. Shallowly erupted Vailulu'u lavas are once again at the lower limit of this range, representing the greatest losses of water by both open- and closed-system degassing.

Original CO_2 concentrations in the primary magmas can be estimated using the observation that CO_2/Nb ratios are roughly constant in volatile under-saturated MORB melt inclusions from the Siquieros Transform (Saal et al., 2002) – it is still highly uncertain whether or not this ratio applies to OIBs. Using fractionation-corrected Nb concentrations and CO_2/Nb equal to 239 (Saal et al., 2002), 95% of the data suggest primary CO_2 contents in the narrow range of 0.8 to 1.2 wt%.

We have used the H_2O - CO_2 degassing program by Newman and Lowenstern (2002) to estimate the maximum water loss from deep, open-system degassing, and find that a primary magma with 47 wt% SiO_2 and initial volatile content of 1.2 wt% H_2O and 1.0 wt% CO_2 will still have 1.15 wt% H_2O after CO_2 has degassed to ~10 ppm. This difference is within the analytical error for water and evidence that the observed water contents for all lavas except those from the shallow Vailulu'u dredges (63, 68, 70 and 73) are robust estimates of original abundances of water.

At the pMELTS-modeled fractionation pressures of 3-4 kbar, the data indicate that CO₂ is the only strongly oversaturated volatile component, as it is very insoluble in basaltic magmas (Stolper and Holloway, 1988; Dixon et al., 1995; Dixon, 1997). Carbonitite

metasomatism has been documented in Samoan xenoliths (Hauri et al., 1993), and is further evidence for deep CO₂ degassing.

4.4. Fluorine

Raw F contents range from 800 to 1890 ppm and are well correlated with incompatible trace elements including P, Na, K and Ti (fig. 5; Table 1). Fractionation-corrected values for F are 480 to 880 ppm and closely overlap primitive Hawaiian melt inclusions that have 300 to 1000 ppm F (Hauri, 2002). These values are similar to other plume related lavas, but are significantly higher than in MORBs (e.g. Schilling et al., 1980; fig. 5).

There are no resolvable differences in the relative enrichment of F between the different volcanoes or the different dredges; instead there is a constant F/Nd ratio of 21.7±2.6 (2σ) across the whole sample suite, indicating F compatibility equal to that of Nd. Figure 5 shows that Samoa glasses, together with normal MORBs (PETBD database: http://beta.www.petdb.org/), very depleted MORBs (from the Siquieros Transform; Saal et al., 2002; Saal et al., unpubl.) and Primitive Upper Mantle (McDonough and Sun, 1995), all fall along a F/Nd line of 21. Siquieros samples alone (Saal et al., 2002) show a mean F/Nd of 19.4±2.6 (1σ), whereas all MORBs show F/Nd = 20.1±5.8 (1σ). Continental crust is also estimated to have a F/Nd ratio within this range (at 20.7; Wanke et al., 1984). To our knowledge, the only basalts that fall dramatically off this line are melt inclusions from the Austral Islands showing F/Nd ratios that are generally 60-70 (Lassiter et al., 2002). Elevated F may be a characteristic of the HIMU mantle end-member that is represented by lavas from the Austral island chain (Hauri and Hart, 1993). However, Vailulu'u glasses, which are thought to contain a small HIMU component (Workman et al., 2004), show no systematic elevation of F.

4.5. Sulfur

Sulfur in the Samoan glasses ranges from 781 – 2651 ppm, consistent with other oceanic basalts (Wallace and Carmichael, 1992). As discussed above, most glasses are saturated with a sulfide phase, so have likely lost some S by sulfide fractionation. Figure 6 shows the rough positive correlation between S/Dy and Mg# of the whole suite of Samoan glasses (note that S and Dy have been suggested to have similar compatibilities during mantle melting [Saal et al., 2002], so S/Dy should not change as a function of melting or olivine/clinopyroxene crystallization, but will decrease if sulfides are fractionated from a

melt). However, for a given Mg#, there is a very wide range of S/Dy, suggesting that fractionation of sulfides may not be the only factor in controlling relative S depletions.

At the FeO contents of Samoan glasses (10.5 - 14.2 wt%), S concentrations are higher than they are in sulfide-saturated MORBs, possibly because Samoan lavas have higher oxygen fugacities (see Dixon, 1997; Dixon and Clague, 2001). Sulfide saturation is a complicated function of fO_2 , temperature, pressure and melt composition (e.g. Wallace and Carmichael, 1992), making further analysis of S beyond the scope of this paper.

4.6. Chorine

Raw Cl contents of the Samoan glasses fall within the range of 451-1815 ppm (Table 1); fractionation-corrected values are 269-1412 ppm, similar to other plume related magmas (e.g. Dixon and Clague, 2001; Simons et al., 2002), but much greater than estimated parental MORBs (<100; Michael and Cornell, 1998). Glasses from Vailulu'u and dredge 76 from Malumalu have the highest Cl, while Ta'u glasses fill the lower half of the whole range. Cl saturation in basaltic melts is greater than 2 wt% at these water contents (Webster et al., 1999), so we expect that there was no Cl degassing at any point in the evolution of the magmas. Curiously, of all the volatile and trace elements, Cl is best correlated with S across the whole Samoan suite (fig. 7). There is no reason to expect these two elements to have either similar compatibilities or similar behavior in magmatic systems.

Cl concentrations are roughly correlated with other incompatible elements for certain groups of dredges. Figure 8 shows one example of this with the relationship between Cl and K (the element most often used in assessing relative Cl enrichments; see below). The lowest Cl/K ratios (~0.04) are found primarily in glasses from Ta'u and Malumalu (dredge 78), and the highest ratios are from a collection of Vailulu'u dredges that define a Cl/K line of approximately 0.12. Two dredges, 73 and 76, fall "across-trend" in between these two groups of Cl/K. It is important to note that this layout is not unique to Cl and K concentration space, and instead looks very similar for Cl with almost all trace elements.

There is much precedent for interpreting high Cl concentrations in oceanic basalts as being the product of seawater assimilation (Michael and Cornell, 1998; Kent et al., 1999; Lassiter et al., 2002; Simons et al., 2002; Stroncik and Haase, 2004) due to very high concentrations of Cl in seawater and especially in brines (Kent et al., 1999). A study by Michael and Cornell (1998) found that Cl/K ratios in MORBs that are unaffected by seawater contamination range from 0.01 – 0.07, with depleted MORBs at the low end and enriched MORBs at the high end of this range. Samoan glasses overlap with the MORB

range and extend to higher Cl/K (up to 0.15; fig. 8). The question is: how much of this Samoan signal is primary and how much is caused by assimilation? We address this question in the paragraphs below.

By omitting samples that fall along brine assimilation trends, Stroncik and Haase (2004) recently found Cl/K ratios in OIBs to be a function of source signatures, as recognized by their correlation with Sr and Pb isotopic values. Enriched mantle (EM1 and EM2) lavas have low Cl/K ratios (down to 0.02) and HIMU lavas have Cl/K ratios up to 0.08 (Stroncik and Haase, 2004). Also, plume-influenced glasses from the Easter Seamount Chain, having Cl concentrations interpreted as primary based on good correlations between Cl and La, show Cl/K ratios from 0.05 up to 0.12 (Simons et al., 2002), similar to our Samoan glasses.

The relationship between Cl/K and ⁸⁷Sr/⁸⁶Sr for Samoa is displayed in Figure 8. The most extreme EM2 lavas (Malumalu, dredge 78) at high ⁸⁷Sr/⁸⁶Sr have some of the lowest Cl/K. With decreasing ⁸⁷Sr/⁸⁶Sr, Cl/K ratios increase up to a maximum value of 0.15 in the Vailulu'u lavas (that have a small HIMU component; see background section above) and then drop to MORB-type values in Ta'u. This result is in general agreement with Stroncik and Haase (2004), despite our absolute Cl/K and ⁸⁷Sr/⁸⁶Sr ranging to significantly higher values.

The minimum Cl/K ratio from each volcano is ~0.04 and could be taken as the primary composition for all of Samoan magmas before Cl contamination. However, enumerated below are reasons to believe that the Cl signal in Samoa is largely primary, with scatter caused by minimal assimilation:

- 1. There are correlations between high Cl and other trace elements that are not enriched in brines (for example, Cl and Nb show a similar relationship as do Cl and K in fig. 7). A 15% brine derived from seawater contains about 10 wt% Cl (Kent et al., 1999), and it would take only 1% of this brine to elevate a magma's Cl content from 500 to 1500 ppm (almost the full range in Samoa). With the same amount of brine, there would be a negligible effect on most other trace elements (including K), yet we observe increased trace element concentrations with increased Cl. Also, Kent et al. (1999) claim that assimilated material is high in B as well as Cl, but B is not elevated in high Cl Samoan glasses; in fact the highest B concentrations occur is glasses from dredge 78, which have some of the lowest Cl and lowest Cl/K ratios.
- 2. Michael and Cornell (1998) clearly demonstrate that high levels of assimilation occur in lavas that are highly evolved. The opposite is observed for the Samoan glasses: the least evolved Vailulu'u samples (see fig. 2) show the highest Cl and Cl/K ratios (fig. 8).

- 3. The few melt inclusions we have analyzed (Table 2) have Cl contents usually within error of their host matrix glasses. However, this evidence alone may not prove the Cl is primary, since it is possible that Cl in the inclusions was assimilated at the depth of crystal fractionation.
- 4. There is a correlation between Cl/K and ⁸⁷Sr/⁸⁶Sr (fig. 8), which in general mimics the observation of Stroncik and Haase (2004) that EM sources and depleted mantle have low Cl/K while HIMU has higher Cl/K. However, Vailulu'u is in no way end-member HIMU, but instead is EM2 with a small component of HIMU.
- 5. Figure 7 indicates that high Cl magmas also have high S, and no known assimilant contains Cl and S in the required 1:1 ratio. Either high Cl is a primary feature of the lavas, or the correlation between Cl and S is coincidental.

A study on chlorine isotopes (e.g. Magenheim et al., 1995) might provide compelling evidence against an assimilation origin for the high Cl, and ultimately may be the only way to define the proportion of igneous- to seawater-derived Cl.

5. Source Variations In Water Enrichment

Since CO₂ is severely degassed, S is subject to sulfide fractionation, Cl is sensitive to seawater contamination, and F abundances do not vary relative to trace element abundances, we focus our attention here on the source variations of water.

During mantle melting, the compatibility of water is similar to Ce and La, as determined both observationally (e.g. Michael, 1995; Danyushevsky et al., 2000) and experimentally (Hauri et al., in revision). As such, ratios of H₂O/Ce and/or H₂O/La have been used as direct indicators of source compositions and the extent to which water is enriched or depleted. For example, low H₂O/Ce ratios in Loihi magmas (Dixon and Clague, 2001) and plume-influenced MORBs (Dixon et al., 2002) have been used to argue for the presence of recycled oceanic crust and sediment that had been efficiently dehydrated during subduction.

The relationship between H_2O/La and $^{87}Sr/^{86}Sr$ in Samoan basalts (fig. 9) extends the observation by Dixon et al. (2002) that enriched mantle sources have relative depletion of water (note that H_2O/La is used rather than H_2O/Ce since H_2O and La are most similar in compatibility during melting of garnet lherzolite [Hauri et al., in revision]). The Samoan glasses have a wide range in $^{87}Sr/^{86}Sr$ (0.7045 – 0.7089) that is inversely correlated with H_2O/La (ranging from 120 to 316). The highest H_2O/La ratios in Samoa overlap with the low H_2O/La end of the plume-influenced Discovery lavas from the S. Atlantic (Dixon et al.,

2002). Otherwise, all Samoa glasses have much lower H_2O/La ratios than do average MORBs or average Loihi lavas (Dixon and Clague, 2001; Dixon et al., 2002). Ultimately, it seems that the greater the "enrichment" of the mantle (in terms of heavy isotopes and trace elements), the greater the apparent dehydration.

We do not believe that this relationship is an artifact of secondary processes in Samoan glasses. There is no coincidental correlation between ⁸⁷Sr/⁸⁶Sr and dredge depth (i.e. possible degassing). In the above section on water and carbon dioxide, we have shown that degassing has not resulted in significant water loss for any of the samples plotted in Figure 9. As supporting evidence, water contents in melt inclusions are equal to or less than the water contents in the glasses at equal or higher CO₂ (Tables 1 and 2). The slightly higher water in some glasses may reflect a greater degree of fractionation of these melts – this is not confirmed since we lack major element compositions for the melt inclusions. For the sake of argument, if the water content in samples from dredge 78 was the same as the highest water in Samoa, then the H₂O/La ratio would be ~200 (instead of 120) – still well below values for MORB, Loihi and Discovery. Finally, if water has been assimilated from seawater or brine, then the H₂O/La ratios plotted in Figure 9 would be maximum values, and the signal we observe would be a dampened one; however, H₂O/La does not correlate with any potential "assimilation" proxies such as Cl/K.

6. Origin Of EM2's "Dehydration"

The origin of the EM2 mantle end-member is classically explained as the recycling of oceanic crust plus terrigeneous sediment through subduction zones (e.g. Weaver, 1991). However, our recent work on extreme EM2 lavas from Samoa (found at Malumalu, in particular) argues against simple mixing of a sedimentary component into depleted mantle based on trace element and isotopic trends in the Samoan lavas (see Workman et al. [2004] for detail). Instead, the origin of EM2 is explained as ancient (2.5 Ga), recycled, metasomatized lower oceanic lithosphere; a small degree, upper mantle melt (like those imagined to be formed at depth beneath mid-ocean ridges or island arcs) impregnated a depleted lithosphere, then the material was subducted and stored in the mantle (Workman et al., 2004). This is simply one scenario that works to explain the observations in Samoa that the EM2 source has a trace element pattern more reflective of upper mantle magmatic fractionation processes than sedimentary compositions, but does not completely rule out a

subduction-related origin for EM2 (in other words, we do not necessarily know *where* the metasomatism has taken place).

The above model for the *formation* of EM2 cannot account for the low H₂O/La ratios observed in the Samoan glasses since (non-arc) mantle melting cannot produce large fractionations of water from La owing to their similar compatibilities. In addition, since the model EM2 reservoir is created in the lower oceanic lithosphere, it will be isolated from the subduction zone dehydration processes affecting the crust/sediment package in the upper part of the subducted slab. However, the *aging* of EM2 *can* account for our observations. This is how: very fast diffusion of hydrogen in olivine (Mackwell and Kohlstedt, 1990) will cause recycled material with high initial water content to lose water to the drier ambient mantle during storage within the deep Earth.

Mackwell and Kohlstedt (1990) provide an Arrhenius equation for the temperature dependence of hydrogen diffusion in olivine; this leads to a diffusion coefficient of 8.9×10 ⁹ m²/s at 1500°C, for the fast [100] direction. Using a constant temperature of 1500°C, in 2.5 Ga (the estimated age of EM2), a 26 km thick slab can be fully equilibrated with the ambient mantle using the approximation that equilibrated distance, $x = \sqrt{\text{diffusion}}$ coefficient x time). At an assumed upwelling rate of 1 cm/yr, water abundance can be fully equilibrated over 4 km with the surrounding mantle during convective transport through the upper mantle (660 km) alone. Since much of the mantle is at temperatures higher than 1500°C, these are conservative estimates; the effects of enhanced diffusion through grain boundaries will also serve to aid equilibration. It is unknown whether or not similar diffusion rates for hydrogen apply to other major mantle minerals. If helium can be used as an analogue for H, at 1500°C, He diffusion in clinopyroxene and garnet is equal or faster than that in olivine (Trull et al., 1991; Dunai and Roselieb, 1996). In contrast to water, most other elements will have much smaller equilibration distances; for example, rare-earthelement diffusion rates at the same temperature results in the full equilibration of less than 10 meters in 4 billion years (Hofmann and Hart, 1978; Van Orman et al., 2001). As a result, the enhanced solid-state mobility of hydrogen in the mantle can potentially shift the H₂O/La ratio in the absence of any melting or mantle-fluid interactions.

Figure 10 is a conceptual diagram for the diffusive equilibration of water during the recycling and storage of EM2. Workman et al. (2004) give an estimate for the trace element composition of EM2: La = 0.895, Ce = 1.923, and if H_2O/Ce was initially like the common mantle, FOZO (i.e. 200; Dixon et al., 2002), then the starting H_2O content of EM2 would be 385 ppm. The Depleted MORB Mantle (DMM) is estimated to contain 0.19 ppm La and

0.55 ppm Ce (Workman and Hart, 2005). At an H_2O/Ce ratio of 150 (Dixon et al., 2002), H_2O in DMM = 82.5 ppm, less than 25% that of EM2. Assuming all reservoirs here are peridotite lithologies, when "wet" EM2 is introduced to "dry" depleted mantle, water will diffuse out of EM2 and into the "infinite sink" of ambient depleted mantle until their water activites are equal (at DMM = 82.5 ppm). When EM2 equilibrates with the depleted mantle and the two components have the same water content, the resulting H_2O/Ce is 43 and H_2O/La is 92 in the EM2 component; these values are nearly identical to the values seen in the high $^{87}Sr/^{86}Sr$ glasses from Malumalu (see fig. 9 for H_2O/La). Less-enriched sources, lying initially between EM2 and DMM along a H_2O/La = 500 line, will lose water when recycled to the mantle until they reach the 82.5 ppm water line. Thus, in aged sources, H_2O/La increases as the degree of enrichment decreases. This analysis shows that the observed differences in H_2O/La ratios in these mantle sources are not the result of different water contents, but instead are due to differences in REE enrichment at similar water contents.

The above discussion is focused on *dehydration* of enriched reservoirs, but the same thinking will apply to *re-hydration* of dry mantle reservoirs. If subducted material becomes extensively dehydrated (e.g. as called upon by Dixon et al. [2002]), storage within the mantle will result in a *re-hydration* of the slab until the slab and mantle are in chemical equilibrium in terms of water content. This is a nearly unavoidable consequence of recycling, unless slab temperatures stay very cold (much less than 1500°C) or recycling times are very rapid (< 50 million years).

7. Discussion

Volatile and trace element abundances for nearly 100 submarine basaltic glasses from the three youngest Samoan volcanoes have been presented here. We have found that water abundance relative to trace element abundance correlates inversely with source enrichment. This leads us to believe that, because of very fast diffusion of hydrogen in olivine (Mackwell and Kohlstedt, 1990), recycled material with high initial water content will lose water to the drier ambient mantle during storage within the Earth. However, this is a somewhat incomplete picture since 1) the greatest budget of water in an upper mantle peridotite is not contained in olivine, but in clinopyroxene and orthopyroxene (Hirth and Kohlstedt, 1996; Hauri et al., in revision) and 2) other lithologies (especially pyroxenites,

with little olivine) may be present in the mantle as subducted oceanic crust (e.g. Hirschmann and Stolper, 1996).

The water content of EM2 is estimated to be close to that of the upper mantle (\sim 100 ppm), and differences in the H₂O/La ratio of Samoan mantle components are due to local diffusive homogenization of their water contents during long-term storage in the mantle. Therefore, the differences in water concentrations between Samoan lavas and MORBs are more a difference in degree of melting than source composition. La, H₂O and Ce are enriched in Samoan lavas about 60-fold over their estimated EM2 source concentrations, requiring a degree of melting on the order of 1% (with D_{Ce, water} = 0.01 and aggregated fractional melting). This contrasts with an estimated degree of melting at 6-7% to generate MORB liquids from the upper mantle (Workman and Hart, 2005).

If the diffusion rates of hydrogen in other mantle minerals are similar to olivine, water may be one of the only elements whose abundance is nearly constant over great distances in the mantle, assuming similar source lithologies. This "diffusive dehydration" model has important implications for the interpretation of mantle H₂O/La and H₂O/Ce ratios. Whether or not subducted materials have higher or lower H₂O/La than the upper mantle, slabs and mantle wedge material stored in the deep Earth for significant periods of time will experience temperature-dependent diffusive mobility of hydrogen and will likely lose hydrogen to the ambient mantle.

The direction of diffusive hydrogen transport will depend not on the H₂O/La ratio but on the bulk H₂O contents of juxtaposed mantle sources. Indeed, subducted oceanic crust and sediment could enter the deep mantle with a low H₂O/La ratio but a bulk H₂O content that is still higher than the surrounding mantle, and thereby lose even more water via "diffusive dehydration". Alternatively, if dehydration and melting are extremely efficient in subduction zones, then recycled material may enter the deep mantle extremely dry, and become re-hydrated via diffusive exchange with the surrounding mantle. Either way, an evaluation of the absolute abundance of water in mantle sources requires more than simply knowing the ratio of H₂O to REE in derivative lavas. Robust estimates of lithophile element concentrations, from accurate inversion of complete major element, trace element and isotopic data, must be the starting point for investigating the variability of water in the mantle in any tectonic setting.

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Figures

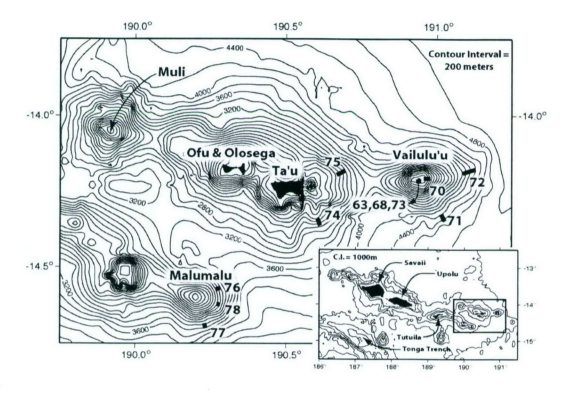


Figure 4-1. Location map for Samoan glasses. Thick lines with adjacent numbers mark individual dredges; note that some dredges traverse more contours than others. Depth contours are plotted using AVON 2/3 shipboard bathymetry data. Inset shows greater Samoan region, including the northernmost extent of the Tonga Trench.

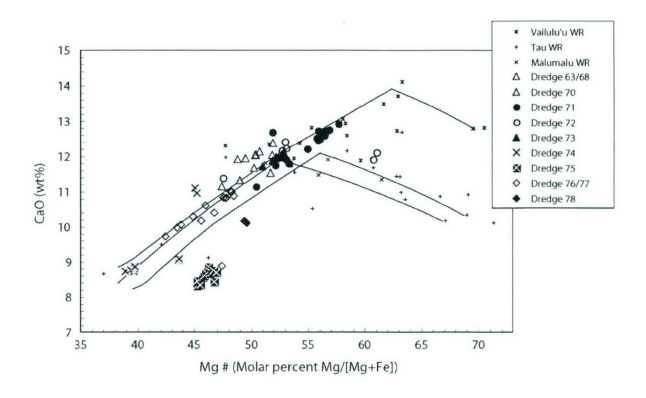


Figure 4-2. CaO plotted with Mg# for Samoan whole rocks (Workman et al., 2004) and glasses (this study). Crystal fractionation trends plotted here have been calculated using pMELTS predicted liquid lines of decent (Ghiorso et al., 2002), starting with some of the most primitive whole rock compositions. Best fits to the observed glass compositional trends are from pMELTS anhydrous runs at pressures of 3-4 kbar. Runs with 1 wt% water are nearly identical to those plotted here, but with 1 kbar higher pressure.

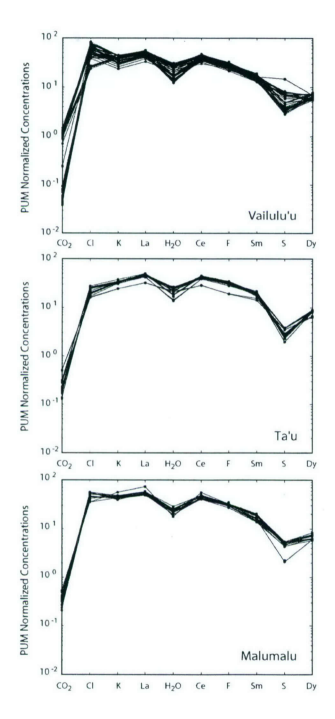


Figure 4-3. Fractionationcorrected trace and volatile element concentrations, listed in the order of increasing compatibility during mantle melting. Values normalized to Primitive Upper Mantle (PUM) of McDonough and Sun (1995), except for water (at 335 ppm assuming $H_2O/Ce = 200$) and carbon dioxide (at 157 ppm assuming $CO_2/Nb = 239$ [Saal et al., 2002]).

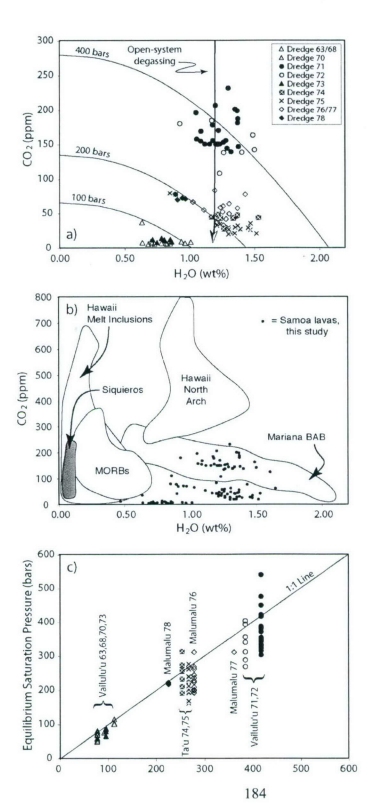


Figure 4-4. Panel a) shows water and carbon dioxide contents of Samoan glasses plotted with equilibrium vapor saturation curves and degassing trends of Dixon (1997);because opensystem degassing trends are nearly vertical on this diagram, we expect that water loss has negligible for all glasses except those at pressures less than 100 bars. Panel b) compares H2O and CO2 contents of Samoan glasses from this study to magmas in other settings. Panel c) plots average dredge for individual depths dredges with calculated saturation pressures (Dixon, 1997) for each sample; most lavas lie near the 1:1 line, so 1) are saturated with an H2O-CO2 vapor and 2) did not flow downhill from shallower extrusion depths. Each dredge traverses multiple depth contours (up to 800 meters; see fig. 1), so there is some ambiguity about the exact collection depth.

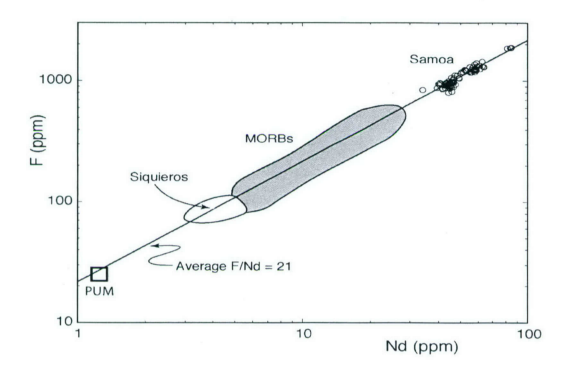


Figure 4-5. Plot of Nd and F concentrations in Samoan glasses from this study, normal MORBs (from the PETBD online database: http://beta.www.petdb.org/), very depleted MORBs (Saal et al., 2002; Saal et al., unpubl.) and Primitive Upper Mantle (McDonough and Sun, 1995). Collectively these oceanic lavas define a constant F/Nd ratio of 21, showing that F has a compatibility equal to Nd during upper mantle melting.

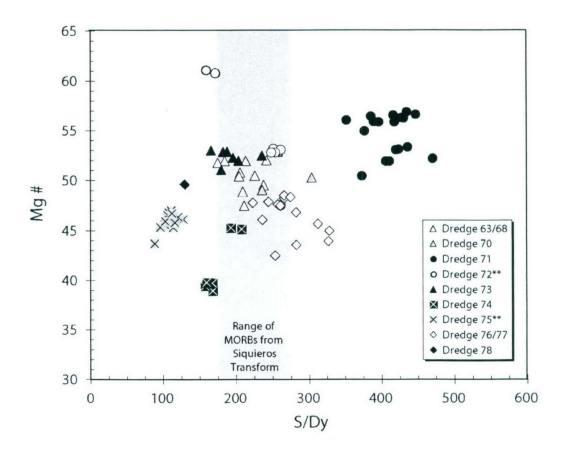


Figure 4-6. Plot of S/Dy with Mg# of the Samoan glasses. Mg# = molar percent Mg/(Mg+0.85Fe). S has been suggested to have similar compatibility to Dy, so S/Dy ratios of melts should be similar to that of their source as long there has been no sulfide fractionation or SO₂ degassing. Since sulfides were observed in all dredges except those marked with **, and there is a rough positive correlation between S/Dy (as well any S/REE ratio) and Mg#, we interpret these lavas to have lost some of their initial sulfur content by sulfide fractionation.

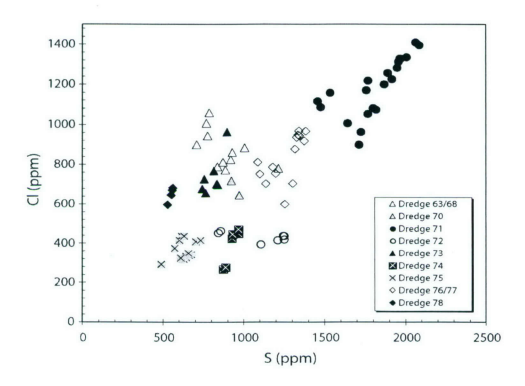


Figure 4-7. Fractionation-corrected S and Cl concentrations in Samoan basaltic glasses. Note from Figure 2 that the glasses with highest S and Cl are from the least evolved glasses of the Samoan suite. Although most glasses are saturated with sulfides, the correlation between Cl and S suggests that S loss by fractionation of a Fe-S liquid may be minimal.

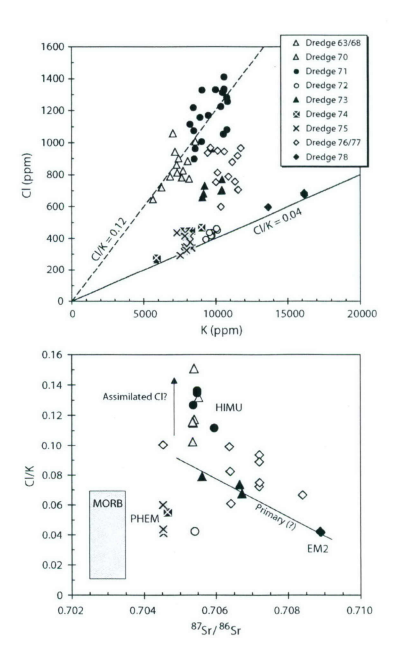


Figure 4-8. Two-panel diagram to show the systematics of Cl in Samoan glasses. Top panel shows fractionation-corrected concentrations of K and Cl; Vailulu'u lavas define a high Cl/K sub-suite, while Ta'u and dredge 78 from Malumalu define a low Cl/K array. The lower panel shows that Cl/K ratios vary with the ⁸⁷Sr/⁸⁶Sr source proxy.

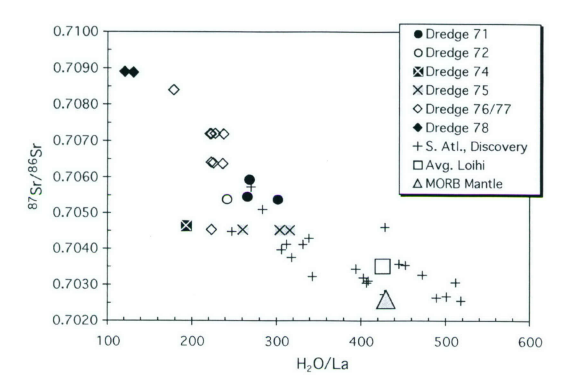


Figure 4-9. Plot of H_2O/La against $^{87}Sr/^{86}Sr$ in Samoan submarine glasses from this study along with plume-influenced glasses from the Discovery anomaly in the S. Atlantic (Dixon et al., 2002), average Loihi glasses (Dixon and Clague, 2001), and average normal MORB based on an H_2O/Ce ratio of 150 (Dixon et al., 2002). Note that the shallowly dredged, degassed Vailulu'u samples (from dredges 63, 68, 70 and 73) are not included on this plot (see fig. 4). The negative slope here indicates that increasingly enriched mantle has decreasing relative enrichment of water in its source. This water depletion in EM2 is interpreted to be the result of diffusive loss of water during storage of EM2 in a dry ambient mantle. The amount of water lost from EM2 is estimated to be ~75% its original (i.e. "pre-recycled") abundance.

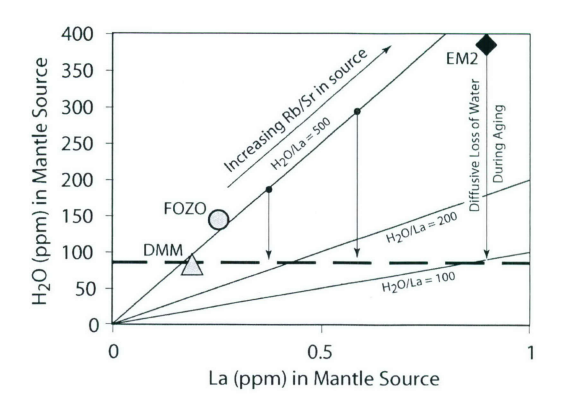


Figure 4-10. This plot is a conceptual diagram for the diffusive equilibration of water during the recycling and storage of EM2. At its formation, EM2 had "normal" enrichment of water, with H₂O/Ce of 200 and H₂O/La of 430 at 385 ppm H₂O. During storage of this material with high H₂O in an ambient mantle with lower H₂O, water is diffusively lost from EM2 – but La is not due to their contrasting rates of diffusion. When EM2 has equilibrated with the upper mantle (82.5 ppm, heavy dashed line), this otherwise enriched reservoir has an H₂O/La ratio of 92. Less enriched sources (that lay in-between EM2 and DMM), will likewise lose water while stored in a drier mantle, but, as a result of lower REE abundance, will equilibrate to higher H₂O/La. This leads to the negative relationship between H₂O/La and ⁸⁷Sr/⁸⁶Sr seen in Figure 9.

Table 4-1. Chemical data for Samoan Glasses, including Major Elements and Fractionation correction factors. Tables

	of minn incurrence			1	0	٧.	orements.	nun	ractionation corre	חוו בחווב	cuon)	actors.	
Sample Volcano	63-13 Vailulu'u	68-03 Vailulu'u	68-03 Rpt Vailulu'u	68-34 Vailulu'u	68-35 Vailulu'u	68-36 Vailulu'u	68-37 Vailulu'u	68-38 Vailulu'u	68-43 Vailulu'u	68-44 Vailulu'u	70-01 Vaihihi'u	70-02 Vaihilii'ii	70-09
Sulfides " Vesicles h	>-		>-	>-	>4	c.4	>-4	>4	C 1	>1	×-	- C	×-
Volatile Content					-	=	=	=		=		-	-
H ₂ O (wt%)	0.83	0.63	0.70	0.82	0.73	89.0	0.85	0.78	0.71	0.64	0 97	101	0.94
CO ₂ (ppm)	9	37	6	7	9	9	7	7	7	· 00	1	0.0	0
F (ppm)	1113	929	924	606	968	884	936	876	915	889	843	935	855
S (ppm)	1318	1127	1153	1472	1353	1322	1469	1044	1410	1361	1387	1876	1484
CI (ppm)	1603	1478	1553	1295	1278	1235	1364	1329	1267	1189	922	1207	1158
# Analyses	2	-	2	3	3	3	3	3	n		9		3,0
Equil. Press. (bars) c	77	69	69	77	09	53	82	71	59	49	104	116	102
Dredge Depth (bar) d	92	78	78	78	78	78	78	78	78	7.8	113	113	113
Major Elements (wt%)8													
Mg# "	47.4	51.9	51.9	52.0	48.8	8.08	49.5	51.8	50.4	50.4	52.8	50.3	49.0
Liquid mass 1	58.9	68.2	68.2	68.3	819	65.8	63.2	8.19	65.1	65.0	70.1	64.8	62.2
Trace Elements (ppm)													
, Ci	8.6	5.9	7.0	5.8	0.9	5.8	6.4	0.9	0.9	5.8	7.0	5.6	6.2
В	3.9	2.7	3.6	2.9	3.0	2.8	3.3	2.9	3.0	2.7	3.7	29	3.5
Ь	2798	1729	2336	1911	9661	1943	2065	1813	1989	1843	2257	1908	2020
×	12153	12597	10298	11785	11034	11126	11509	10927	11877	12551	8030	11785	10023
Ti	20607	20450	19118	21234	21782	21547	20999	19980	21312	21782	17786	21156	20372
Rb	23	30	19	26	25	24	24	25	26	30	14	27	21
Sr	459	582	478	564	563	565	559	510	583	619	411	552	444
\	30	29	26	29	30	29	29	28	29	31	25	30	30
Zr	283	271	253	270	283	278	271	266	280	285	225	272	265
NP	58	56	50	53	54	54	54	54	55	57	39	56	51
Ba	280	304	243	288	278	278	282	255	295	315	171	280	232
La	47	43	39	41	42	42	41	41	42	44	31	43	39
o Ce	100	94	87	91	94	94	06	16	93	76	72	94	87
PN	47.2	43.8	40.5	43.5	44.5	45.6	42.9	42.8	44.6	46.5	34.1	44.1	41.7
Sm	9.1	8.5	8.5	0.6	9.6	9.8	8.7	8.2	8.8	9.1	7.4	8.1	8.3
Eu	2.8	3.1	2.4	3.0	3.0	3.0	3.1	2.9	2.9	3.3	2.1	2.9	2.7
PS	7.0	∞. ∞	6.5	8.1	8.3	8.4	8.0	8.1	8.4	9.3	6.1	8.3	7.4
Dy	6.2	6.1	5.4	6.1	6.4	6.4	6.2	0.9	6.2	9.9	5.4	6.2	6.3
Yb	2.5	2.6	2.0	2.4	2.5	2.5	2.1	2.7	2.5	2.5	1.7	2.7	2.6
Hf	6.9	5.9	6.1	0.9	6.5	6.5	6.1	6.2	6.1	6.2	4.9	5.8	6.2
Th	8.1	5.7	6.5	8.5	5.8	5.9	6.1	5.8	0.9	5.7	5.5	6.1	6.1
U - 26	1.8	1.3	1.3	4.1	1.4	1.3	1.4	1.3	1.3	1.3	1.1	1.3	1.4
S./Sr,	0.70552	0.705396	0.705396								0.705371	0.705359	0.705352

* Presence of sulfides in glass chips; y = observed, n = not observed.

* Vesicle abundance in glass chips; J = Iow (0.5 vol%), m = moderate (5.10 vol%), h = high (10.30 vol%)

vol%).

^c Equilibrium vapor saturation pressure calculated following Dixon (1997).

^d Average dredge depth for each dredge; 10 meters = 1 bar.

^e Sr Isotope analyses on whole rocks previously published by Workman et al. (2004) are in plain text, new analyses on glasses are in bold.

⁶ Repeat analyses are from different glass chips of the same sample.

⁸ Major Elements previously published in Workman et al. (2004).

⁸ Major Elements previously published in Workman et al. (2004).

⁹ Liquid mass = percent of liquid mass remaining after fractionation of soilds from a Mg# 73 initial composition as predicted by pMELTS (Ghiorso et al., 2002). Multiply trace and volatile elements by this number/100 to get fractionation corrected compositions. Amount of solid fractionation = (100 - Liquid remaining).

Sample	71-02	71-03	71-04	71-05	71-06	71-06Rpt	71-09	71-10	71-10Rpt	71-11	71-13	71-13D	71-14
Volcano	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailuluu	Vailulu'u	vailulu'u	Vailulu u	Vailulu
Sulfides ³ Vesicles ^b	>-	>-	>-	>-	>-	>-	>-	>-	>-	>-	>-	>-	>-
Volatile Content													
H2O (wt%)	1.18	1.22	1.09	1.06	1.28	1.37	1.18	1.26	1.37	1.23	1.12	1.37	1.22
CO. (nnm)	179	171	156	158	156	188	156	154	182	151	151	199	152
E (npm)	003	080	037	865	070	1064	947	996	1026	915	878	796	949
(indd) I	500	2000	150	500	717	1001	2500	25.75	0701	0736	2441	2000	7537
S (ppm)	2537	7197	74/4	0907	4/67	7404	2208	7/67	2308	2007	1447	2400	1007
CI (ppm)	1490	1752	1651	1558	1/18	14/8	1656	1/44	1610	7751	1300	1971	0/1
# Analyses	3	3	n	2	3	4	m	3	4	n	m	n	3
Equil. Press. (bars) c		356	304	333	382	451	323	352	418	355	335	477	332
Dredge Depth (bar) d	417	417	417	417	417	417	417	417	417	417	417	417	417
njor Elements(wt%)8													
Mg# h		57.7	55.9	55.0	51.9	51.9	56.6	55.9	55.9	53.1	53.1	53.3	56.3
Liquid mass		7.67	76.3	74.4	68.2	68.2	77.5	76.2	76.2	70.7	9.07	71.1	77.0
Trace Elements(ppm)													
Li	×	63	6.1	6.3	6.5	6.3	6.1	6.2	5.8	8.9	5.7	6.2	5.6
m	3.0	33	33	23	37	33	33	3.6	3.0	3.0	2.9	3.3	2.7
2 0	1560	1735	1733	1771	1883	1779	1733	1777	1750	1724	1741	1756	1641
- 2	13217	14206	11060	13000	13210	13884	13860	13503	11084	12137	11938	13609	1241
Z F	20215	21156	18727	21038	20685	20685	20764	20764	19353	19040	19118	20764	1951
8 P	32	36	200	33	30	34	34	33	29	28	28	32	30
5.	267	209	544	597	268	288	591	585	545	543	545	586	566
; >	28	29	27	29	28	29	28	29	27	27	28	30	29
Zr	271	285	260	286	270	279	278	282	260	263	258	294	264
NP	55	59	53	58	55	58	57	57	53	53	52	57	54
Ba	305	327	287	319	299	314	321	314	284	279	276	312	296
La	44	47	41	46	45	45	45	46	41	39	39	44	41
Ce	96	103	68	102	86	76	86	100	88	98	87	66	92
PN	45.4	47.5	41.8	46.7	45.6	44.9	45.9	46.3	42.4	39.5	40.7	45.5	42.9
Sm	8.2	80.00	8.9	0.6	9.2	8.5	9.4	9.1	8.1	8.4	8.2	8.1	8.3
En	3.2	3.2	2.8	3.2	2.8	3.4	3.2	3.2	2.8	2.8	2.7	3.0	3.0
Cd	0.6	9.3	7.5	9.3	7.9	9.2	9.1	0.6	8.2	8.1	8.3	8.9	8.8
Dv	0.9	6.4	5.5	6.3	5.9	6.0	6.2	6.3	6.1	5.8	5.5	5.9	6.1
X.	2.3	2.5	2.2	2.5	2.4	2.4	2.3	2.5	2.5	2.4	2.7	2.4	2.7
H	5.9	6.1	0.9	6.1	6.4	6.1	5.8	6.2	5.9	5.3	5.8	5.9	0.9
1	8.0	6.4	5.5	6.2	8.9	5.9	0.9	6.5	5.3	5.0	5.4	5.9	5.5
n	1.4	1.4	1.3	1.4	1.5	4.1	1.4	1.4	1.2	1.2	1.2	1.3	1.3
870 860 e	0000000												

Table 4-1, Page 3

Sample	61-17	01-1/	77-11	· desert			64-11	04	17-11	07-11	101107-11	1 de 17 mm	
Volcano	Vailulu'u	Vailulu'u											
Sulfides a	y	y	Y	E	x	^	>	>	>	>	u	-	-
Vesicles b	_	-	-	_	. —	. —	.—		_	_	: -	: =	-
Volatile Content													
$H_2O(wt\%)$	1.38	1.32	1.09	1.30	1.22	1.29	1.15	1.20	1.05	1.20	1.35	1.23	1.26
CO_2 (ppm)	147	140	169	233	151	145	151	151	197	207	202	109	139
F (ppm)	918	944	957	1110	096	983	962	973	066	1056	1077	1188	1060
S (ppm)	2619	2569	2232	2261	2551	2651	2563	4834	2323	2304	2435	1870	1757
CI (ppm)	1545	1775	1715	1670	1726	1818	1711	1754	1400	1377	1570	670	623
# Analyses	3	3	2		r			, ~			4	200	220
Equil. Press. (bars) c		373	388	542	330	344	316	221	346	386	173	070	213
Dredge Depth (bar) d	417	417	417	417	417	417	417	417	417	417	423	384	384
Major Elements(wt%)8												100	100
Mg# h		52.2	50.5	50.5	56.3	56.7	56.9	55.8	56.5	56.1	56.1	47.5	53.1
Liquid mass 1		8.89	65.2	65.2	77.1	7.77	78.2	76.0	77.3	76.6	76.6	59.1	70.6
Trace Elements(ppm)													
Li	5.6	5.8	0.9		7.4	6.3	6.2	6.5	6.3	6.2		7.1	6.2
В	2.7	3.0	3.4		4.0	3.5	3.4	3.6	3.2	3.3		4.0	3.4
Ь	1641	1781	1764		2229	1754	1779	1898	1780	1809		2105	1870
×	12413	12260	12567		11739	13624	13547	13149	13953	13747		15708	13716
Ţ	19510	19353	19040		19824	20764	20764	20999	20842	20921		24760	23663
Rb	30	28	31		24	32	31	32	34	32		37	33
Sr	995	554	575		492	583	582	292	592	165		999	919
~	29	28	27		26	29	29	28	29	29		34	33
Zr	264	261	270		258	284	280	280	278	280		321	302
NP	54	54	55		54	99	58	57	57	57		19	59
Ba	296	287	298		259	313	311	310	321	316		361	322
La	41	40	41		42	46	45	45	46	46		51	47
Ce	92	88	06		06	46	76	86	100	101		112	103
PN	42.9	40.1	40.5		43.2	45.5	43.3	43.5	46.4	46.8		51.4	49.2
Sm	8.3	7.7	8.5		8.9	8.8	7.6	8.8	9.2	9.3		10.9	10.2
Eu	3.0	2.9	3.0		5.6	2.9	3.2	2.9	3.0	3.1		3.7	3.3
PS	8.8	8.4	8.5		8.9	9.8	8.5	8.2	8.8	8.8		10.3	7.6
Dy	6.1	5.5	0.9		0.9	5.9	5.9	6.2	0.9	6.5		7.2	7.0
Yb	2.7	2.4	2.4		2.0	2.5	2.7	2.3	2.4	2.4		2.9	2.7
Ht	0.9	5.6	0.9		6.3	6.1	6.2	6.2	6.2	6.2		7.1	6.9
T	5.5	5.4	5.5		8.9	5.8	5.9	6.4	6.3	6.5		8.9	0.9
U 86- 8	1.3	1.3	1.2		1.5	1.4	1.3	1.4	1.4	1.5		1.5	1.4
2 100/10			0 705172	0 705 473									

Table 4-1, Page 4

Sample Volcano	72-07	200				1000			1 4 1	12 12	72 12	72 15	74-01
Volcano *		72-10	772-17	/2-12Kpt	72-13	73-01	73-03	73-04	73-07	71-5/	(1-0)	13-13	10-1-1
	Vailulu'u	Ta'u											
Sulfides a	п	п	п	п	п	п	У	y	п	п	y	п	
Vesicles b	_	_	_	_	-	31%	20%	٠ ح	ч	٦	.—	۲	-
Volatile Content													
H ₂ O (wt%)	1.49	0.93	1.40	1.50	1.17	0.75	0.71	0.87	0.78	0.81	0.85	0.81	1.26
CO ₂ (ppm)	164	181	139	143	185	∞	13	6	15	12	13	11	28
F (ppm)	1148	867	1050	1109	921	686	866	1022	955	982	696	932	801
S (ppm)	1771	1010	1728	1784	1041	1209	1138	1227	1055	1162	1001	1293	984
CI (ppm)	869	547	969	628	267	1022	1098	1025	596	1097	940	1390	651
# Analyses	3	3	7	5	3	3	3	3	3	3	2	3	3
Equil. Press. (bars) c	405	289	357	394	341	99	99	87	80	80	06	80	
Dredge Depth (bar) d	384	384	384	384	384	96	96	96	96	96	96	96	254
Major Elements(wt%)8													
Mg# h	53.0	61.1	52.8	52.8	8.09	52.3	51.0	52.0	53.0	52.9	52.8	52.5	
Liquid mass 1	70.5	83.0	6.69	6.69	82.0	6.89	66.4	68.3	70.4	70.1	70.0	69.3	
Trace Elements(ppm)													
Li	6.4	6.2	6.3	6.2	5.9	5.5	6.2	5.5	5.7	5.6	5.7	5.8	
В	3.5	3.5	3.6	3.9	3.4	4.0	3.0	4.0	2.7	4.0	2.8	4.5	
Ь	1885	1816	1853	1802	1831	1763	1886	1773	1764	1813	1746	1986	
×	13793	12168	13808	13639	12245	15156	13930	15264	13026	14911	12965	13961	
Ti	23741	19824	23585	23428	20215	19432	20529	19432	19902	19589	19745	19275	
Rb	33	31	32	32	30	37	32	37	32	35	31	30	
Sr	620	559	612	665	570	585	995	589	570	587	267	543	
7	33	29	33	32	30	29	29	29	29	29	28	27	
Zr	300	280	300	297	291	287	281	287	275	283	569	280	
NP	09	55	59	58	99	58	28	58	55	28	55	58	
Ba	324	294	321	314	297	388	325	392	306	379	308	354	
La	47	43	46	45	44	52	45	52	44	52	42	51	
Ce	104	93	101	66	76	109	76	109	96	109	93	105	
PN	50.3	44.9	49.5	48.4	45.4	46.6	44.2	46.5	42.9	45.5	43.6	44.1	
Sm	9.3	8.6	9.4	10.7	9.5	8.4	8.8	0.6	8.8	8.8	8.4	8.0	
Eu	3.4	3.1	3.3	3.4	3.0	3.1	3.3	3.1	3.1	3.1	3.2	2.9	
PS	6.6	8.7	7.6	9.6	9.3	8.9	8.8	8.5	8.1	8.8	8.5	7.3	
Dy	8.9	6.4	8.9	7.2	6.1	6.2	6.3	0.9	6.4	6.2	0.9	5.5	
Yb	2.9	2.4	3.1	2.7	2.5	2.4	2.6	2.5	2.6	2.5	2.5	2.4	
Hf	6.9	5.9	9.9	6.4	6.2	6.3	6.2	6.1	5.8	6.3	5.9	0.9	
Th	5.8	5.7	5.9	5.9	0.9	7.3	6.3	7.5	6.1	7.8	5.9	8.3	
D	1.4	1.3	1.4	1.2	1.3	1.5	1.4	1.5	1.3	1.3	1.3	1.5	
87S1/86Sr c						0.70672	0.705616			0.706653			0.704686

Table 4-1, Page 5

Sample	70-4/	/4-03	74-05	74-10	74-11	74-12	74-13	74-14	75-01	75-02	75-03	75-04	75-05
Volcano	Ta'u												
Sulfides a	n	u	п	п	E	r.	>	>	u	2	-	-	-
Vesicles b	_	-	-	-	-	_	. —	. —	E	ш	E	E	Ε
Volatile Content													
H ₂ O (wt%)	1.36	1.16	1.34	1.37	1.22	1.24	1.33	1.53	1.33	1.52	1.35	1.30	1.39
CO_2 (ppm)	46	45	44	48	49	44	44	44	21	21	21	20	27
F (ppm)	1872	802	1871	1866	1856	818	1854	1887	1339	1371	1235	1338	1262
S (ppm)	2107	1462	2092	2044	2017	1498	2022	2098	866	1034	1132	666	1189
CI (ppm)	982	451	975	066	928	469	971	1023	689	734	657	692	674
# Analyses	4	3	3	3	3	3	3	3	3	3	3	3	3
Equil. Press. (bars) c	263	193	256	275	233	212	258	314	214	271	220	205	240
Dredge Depth (bar) d	254	254	254	254	254	254	254	254	268	268	268	268	268
Major Elements(wt%)8													
Mg# h	38.9	45.3	39.8	39.4	39.7	45.1	39.7	39.8	45.6	45.3	46.3	46.8	46.1
Liquid mass	44.1	59.5	46.1	45.2	45.9	59.2	46.0	46.1	60.3	9.69	62.1	63.2	9.19
Trace Elements(ppm)													
Ľ	10.2	0.9	10.3	10.3		5.9	10.2	10.2	8.4	0.6	8.0	8.8	7.8
В	6.4	3.4	6.5	9.9		3.4	9.9	8.9	5.1	5.8	4.8	5.5	4.7
Ь	4327	2003	4351	4364		1965	4313	4341	3330	3865	3077	3757	3031
K	18191	9854	17961	17900		0066	16720	19509	13655	12229	12766	12475	12720
Ţi	24211	37767	23741	23741		37532	23271	23898	24368	23428	26797	23976	27189
Rb	38	21	38	39		23	36	40	28	24	27	24	27
Sr	664	299	664	671		919	650	649	730	029	735	685	734
7	28	35	59	09		35	57	59	45	43	43	45	44
Zr	287	314	594	604		308	557	585	445	431	429	442	427
NP	98	49	98	88		48	98	88	65	61	19	63	62
Ba	373	209	370	383		213	366	388	287	262	275	271	270
La	69	35	69	71		35	49	71	50	48	47	50	47
Ce	191	81	162	991		80	158	165	116	112	109	117	109
PN	84.3	44.3	84.1	84.5		45.8	81.3	84.9	61.0	58.0	58.4	60.7	58.0
Sm	17.3	7.6	17.6	17.8		9.01	17.1	18.0	12.7	11.7	12.2	13.3	12.5
Eu	5.6	3.3	5.7	5.8		3.2	5.6	5.7	4.5	4.0	4.2	4.2	4.4
PS	17.4	7.6	17.5	18.3		10.0	17.1	18.1	13.2	11.5	12.9	11.5	12.8
Dy	12.5	7.6	12.7	13.0		7.2	12.1	13.1	7.6	0.6	9.3	9.2	9.3
Yb	5.1	3.0	4.9	5.1		2.9	5.1	5.5	3.7	3.5	3.7	3.7	3.7
H	12.7	6.9	12.4	12.8		6.9	12.6	13.4	9.4	7.6	9.2	8.6	9.3
T	8.2	4.2	8.2	7.9		4.0	8.2	8.7	5.7	6.5	5.2	6.7	5.4
D	2.1	6.0	2.1	2.2		1.0	2.2	2.2	1.5	1.6	1.3	1.5	1.4
0 - 70													

Table 4-1, Page 6

Sample	75-07	75-08	75-10	75-11	75-12	75-13	75-14	75-15	75-16	75-17	75-18	76-01	76-02
Volcano	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Malumalu	Malumalu
Sulfides a	r	п	u	п	E	L	r	п	r	п	r	y	y
Vesicles b	ш	ш	ш	ш	ш	ш	ш	ш	ш	ш	ш	-	1
Volatile Content													
$H_2O(wt\%)$	1.21	1.30	1.23	1.35	1.47	0.85	1.37	1.47	1.45	0.90	1.30	1.37	1.30
CO ₂ (ppm)	33	30	35	30	26	80	32	36	29	78	27	58	64
F (ppm)	1291	1240	1234	1245	1448	1299	1216	1311	1219	1294	1175	1222	1220
S (ppm)	1086	1027	1039	1040	096	806	1024	1084	1004	904	696	1923	1991
CI (ppm)	551	534	526	531	629	536	519	574	520	541	514	1283	1361
# Analyses	3	3	3	3	3	3	3	3	3	3	3	3	3
Equil. Press. (bars) c	193	216	202	231	263	162	240	276	262	168	213	268	262
Dredge Depth (bar) d	268	268	268	268	268	268	268	268	268	268	268	279	279
Major Elements(wt%)8													
Mg# h	46.3	46.5	46.3	47.0	45.3	43.7	46.0	45.8	45.9	43.6	46.7	46.0	43.5
Liquid mass 1	62.1	62.6	62.0	63.7	9.65	54.2	61.4	8.09	61.1	54.1	63.0	61.4	55.3
Trace Elements(ppm)													
Li	7.7	7.7	7.8	7.9	8.3	9.8	7.9	8.0	7.6	8.6	7.8	9.2	10.7
В	4.9	4.7	4.7	4.8	4.9	5.4	4.8	5.2	4.8	5.6	4.8	4.3	4.6
Ь	3012	3039	3037	3017	3307	3234	3027	3165	3009	3302	3009	2377	2046
×	12613	12700	12775	13107	13762	13805	12977	13103	12773	13935	12648	17745	18166
Ţi	26879	27134	27455	27338	24666	31122	27424	25222	27040	31020	26915	26750	24204
Rb	28	28	28	29	30	30	28	28	29	30	28	43	42
Sr	737	751	745	740	751	999	739	749	755	650	747	734	658
>	43	44	44	44	46	49	44	44	44	49	43	36	33
Zr	416	424	427	428	458	471	431	428	427	471	421	371	321
NP	09	19	19	62	65	19	19	63	19	99	19	70	74
Ba	569	272	275	278	297	287	274	282	275	286	270	403	402
La	46	47	47	48	52	51	47	48	47	20	47	58	57
Ce	106	=	110	Ξ	120	119	110	113	110	119	108	125	122
PZ	8.99	58.5	58.6	58.9	62.7	63.6	58.8	59.2	57.7	64.2	55.5	57.3	52.8
Sm	12.1	12.2	12.4	14.0	13.8	14.7	12.4	12.8	12.7	14.3	11.6	12.2	10.2
Eu	4.1	4.3	4.3	4.3	4.5	4.9	4.4	4.2	4.2	4.3	4.2	4.2	3.7
PS	12.4	13.2	12.9	13.1	13.5	13.9	13.3	13.1	13.1	13.8	13.1	11.8	10.6
Dy	8.9	9.6	9.3	9.3	6.6	10.2	9.1	9.3	6.7	10.1	9.8	8.1	7.0
Yb	3.6	3.7	3.8	3.8	4.1	4.3	4.0	3.7	3.7	4.2	3.8	2.8	2.8
Hf	0.6	9.1	0.6	9.1	9.5	6.6	8.9	9.3	8.8	10.1	8.6	8.0	8.9
T	5.1	5.3	5.2	5.2	5.6	5.5	5.3	5.4	5.2	5.8	5.0	7.1	7.4
D	1.3	1.3	1.4	1.3	1.5	1.5	1.3	1.4	1.4	1.5	1.3	1.6	1.6
87Sr/86Sr c			0.704533					0.704527				0.707192	0.707202

Table 4-1, Page 7

Sample	/6-03	76-04	76-05	90-92	20-92	80-92	76-10	76-11
Volcano	Malumalu							
Sulfides a	y	×	y	×	Y	L	^	>
Vesicles b		-	. –	. –	. —	_		
Volatile Content								
H ₂ O (wt%)	1.25	1.11	1.20	1.03	1.30	1.21	1.26	1.24
CO ₂ (ppm)	52	55	84	19	50	37	33	59
F (ppm)	1180	1237	1182	1236	1300	1236	1409	1228
S (ppm)	1735	2023	1936	1894	2391	1992	2065	2277
CI (ppm)	1074	1348	925	1195	1725	1403	1541	1615
# Analyses	3	3	3	3	3	6	2	
Equil. Press. (bars) c	228	192	311	194	244	199	209	236
Dredge Depth (bar) d	279	279	279	279	279	279	279	279
Major Elements(wt%)8								
Mg# h	47.7	47.6	47.4	46.8	43.9	48.2	42.5	44.9
Liquid mass 1	65.5	65.1	64.8	63.2	56.1	66.7	52.7	58.7
Trace Elements(ppm)								
r, II	8.9	0.6	9.8	12.8	10.6	10.9	6.6	6.6
В	4.4	4.5	3.9	0.9	4.6	5.3	5.0	4.6
Ь	2369	2499	2059	2605	2214	3200	2730	1982
×	17639	17133	16015	17961	17195	14176	19264	17287
Ţ	26758	26484	25308	23820	24055	24682	25857	23976
Rb	44	41	37	41	39	28	46	42
Sr	728	402	641	809	633	594	783	029
Y	35	35	34	30	33	33	35	33
Zr	365	368	301	310	321	333	380	319
NP	69	70	99	65	72	62	78	71
Ba	402	390	354	386	390	320	439	394
La	99	57	51	58	58	51	19	26
Ce	123	123	112	119	122	116	131	120
PZ	57.9	57.6	50.6	52.2	55.2	56.2	59.0	52.5
Sm	11.2	11.7	10.5	10.5	10.8	12.0	11.5	6.6
Eu	4.1	3.9	3.7	3.1	3.4	3.1	4.4	3.9
PS	11.2	11.1	10.0	8.5	10.2	8.5	11.7	10.8
Dy	7.8	7.8	7.4	6.7	7.3	7.5	8.1	6.9
Yb	2.7	2.8	2.7	2.1	2.5	2.2	3.1	3.0
Hf	8.5	8.1	6.7	7.7	7.2	7.6	8.0	6.9
Th	7.0	7.6	8.9	10.3	8.7	8.3	8.0	7.6
n	1.6	1.7	1.5	1.9	1.7	1.9	1.8	1.7
870 860 e				200000	0.704530	ALC2010		10000

Table 4-1, Page 8

Sample	76-12	76-13	76-14	90-77	78-01	78-02	78-03	78-10
Volcano	Malumalu							
Sulfides a	^	۸	^	^		>		^
Vesicles b	_	_		, _	4	٠, ـ	24%	ر د
Volatile Content								
H-O (wt%)	1 20	1 28	1 26	1.43	0.01	0.05	0.07	080
(41/4)	07:1	27:1	07:1	0,0	0.0	55.5	10.0	10.0
CO ₂ (ppm)	44	74	10	8/	0/	13	7/	9/
F (ppm)	1245	1221	1211	1173	1254	1226	1310	1206
S (ppm)	2089	2053	2225	1947	831	821	838	781
CI (ppm)	1396	1436	1562	1055	1004	896	1021	988
# Analyses		۲	٠	"	~			3
Fouril Proce (hore)	200	224	245	311	,	218)	223
Darder Press (Vars)	202	177	010	361	711	917	311	326
Dredge Deptin (bar)	617	617	617	201	077	077	077	077
Major Elements(wt%)								
Mg# #	47.8	48.5	45.6	48.3		49.4		49.6
Liquid mass 1	65.8	67.3	60.4	6.99		0.79		67.4
Trace Elements(ppm)								
7.1	8.9	8.7	10.1	10.3	11.5		12.1	13.5
8	4.5	4.4	4.5	5.3	6.2		6.3	7.8
۵	2434	2409	1976	2390	2078		2114	2685
×	17532	17440	17578	17287	24137		24106	20214
Ti	26640	26484	23898	25230	21234		20921	19824
Rb	42	42	43	41	61		09	46
Sr	723	720	650	644	723		704	620
*	36	36	32	33	33		32	30
Zr	370	372	318	343	384		371	358
NP	71	70	71	71	78		77	72
Ba	397	398	392	380	481		477	405
La	57	57	56	57	92		74	71
Ce	124	125	119	124	155		151	140
PN	58.1	59.0	53.3	56.1	61.4		63.0	55.5
Sm	11.9	12.4	10.6	11.3	10.7		10.8	9.5
Eu	4.2	3.9	3.5	3.3	3.9		3.9	3.0
PS	11.5	11.8	10.8	8.6	11.5		10.9	8.7
Dy	8.5	7.7	7.1	7.1	7.1		7.2	0.9
Yb	2.7	2.9	2.6	2.8	2.7		2.6	2.2
Hf	8.0	7.7	9.9	8.0	8.1		8.1	7.9
Th	7.4	7.2	7.8	8.4	11.6		11.1	12.6
n	1.7	1.7	1.6	1.7	2.2		2.3	2.3
875r/865re		0.706395	0.707198		0.708901		0.708886	

Table 4-2. Volatile Composition of Olivine-Hosted Melt Inclusions

	68-34 mi	71-03mi	71-03mi	71-06 mi	72-12 mi	72-12 mi
H ₂ O (wt%)	0.88	1.22	0.67	1.21	1.48	1.32
CO ₂ (ppm)	35	164	53	453	509	515
F (ppm)	935	098	879	874	1107	1022
S (ppm)	2028	2265	1979	2242	2348	2220
Cl (ppm)	1296	1598	1256	719	541	517
	76-07 mi	76-08 mi	76-10 mi	76-13 mi	76-14 mi	77-06 mi
H ₂ O (wt%)	1.10	1.14	0.93	1.12	1.08	1.35
CO ₂ (ppm)	95	37	26	388	57	161
F (ppm)	1129	1233	1182	1150	1170	1239
S (ppm)	2195	2019	1661	2121	2033	1952
Cl (ppm)	1403	1411	1279	1295	1470	1051

Chapter 5:

Assessment of recycled, slab-derived material in enriched lavas from Samoa: evidence from oxygen isotopes

Abstract

Oxygen isotope compositions of olivine phenocrysts from Samoan lavas, ranging from δ¹⁸O of 5.1-5.6‰, reconfirm that lavas from Enriched Mantle 2 (EM2) plumes display higher primary δ^{18} O values than any other mid-ocean ridge or hotspot setting. We find that δ¹⁸O correlates well with ⁸⁷Sr/⁸⁶Sr, ²⁰⁸Pb/²⁰⁴Pb and incompatible element ratios such as La/Sm. There is no correlation between δ^{18} O and Mg#'s of whole-rocks or olivines, so we interpret the 818O values to represent primary melt compositions. Clinopyroxene phenocrysts were also analyzed from a few samples and show a similar range in δ^{18} O as the olivines; unlike a typical equilibrium fractionation of ~0.4% for $\delta^{18}O_{CDX} - \delta^{18}O_{OD}$, most clinopyroxenes are lower in δ^{18} O than their coexisting olivines. We test two models for the origin of EM2's enrichment in 18O, incompatible trace elements and Sr-Nd-Pb radiogenic isotopes; both are based almost entirely on comparisons of trace element patterns between a calculated source for the most EM2-rich lavas and our two model outputs. The first model is inspired by the classic interpretation that EM2 represents mantle that has been enriched by recycled marine sediments. The second model is one in which a subduction-zonedelineated mantle wedge is metasomatized by a melt from oceanic crust that has been subducted to eclogite facies pressure-temperature conditions. Each model has significant discrepancies with trace element patterns observed for Samoa, but the metasomatic model overall shows a better fit.

1. Introduction

Oxygen isotopes have become a commonly used proxy for the identification of ancient, recycled crustal components in arcs and mantle plumes and/or as a means to identify recent crustal contamination of mantle-derived igneous rocks (Eiler, 2001). The use of oxygen isotopes as a crustal tracer is made possible by the fact that only materials at or near the Earth's surface show great variability in δ^{18} O as a result of low temperature equilibrium fractionation effects. However, δ^{18} O variations in mantle-derived lavas and their phenocrysts are often small and care must be taken not to interpret secondary processes as mantle source signals.

The range in δ^{18} O of olivines from MORBs and upper mantle peridotites is small (5.1 \pm 0.2%; Mattey, 1994; Chazot et al., 1997; Eiler et al., 1997; Eiler, 2001) and overlaps with values obtained from lunar rocks (Clayton et al., 1971; Clayton et al., 1972, Wiechert et al., 2000). Deviations from these 'primary' mantle values have been found in ocean island basalts (OIBs) and attributed to the addition of sediment/upper-crustal material (in the case of high δ^{18} O) or lower oceanic crust (in the case of low δ^{18} O), either as a present-day assimilant or as a component in the mantle plume (see the review by Eiler, 2001). Most detailed studies of δ^{18} O compositions in OIBs have focused on locations where δ^{18} O is either very similar to MORB (Pitcairn, Hawaii) or is lower than MORB (Hawaii, Iceland, Canaries) (Eiler, 2001). Here we expand on the observation by Eiler et al. (1997) that the EM2 mantle component (Enriched Mantle 2), found most extremely in the Samoan hotspot track and more weakly in the Society Islands, is the only one to consistently show δ^{18} O values that are elevated over MORBs. The Samoan islands and seamounts are now very well characterized in terms of major and trace elements, water content and Sr-Nd-Pb-Hf-Os-He isotopic composition, so there is abundant context in which to put the new oxygen isotope data that is presented in this paper.

Eiler et al. (1997) interpreted the high $\delta^{18}O$ in Samoa as the occurrence of recycled marine sediment in the Samoan plume. Such an explanation for the origin of enrichment in EM2 followed previous suggestions for EM2's origin, but was subsequently refuted by Workman et al. (2004) based on a large suite of trace element and isotopic data. Instead, Workman et al. (2004) called upon metasomatism of the lithospheric mantle (by an upper mantle melt) to explain the formation of EM2; however, their model cannot directly account for elevated $\delta^{18}O$ compositions.

Here we present new δ^{18} O analyses of olivine and clinopyroxene phenocrysts from Samoan lavas and test two models for the generation of EM2. The first is the standard sediment-recycling model. The second expands upon the metasomatic model of Workman et al. (2004) by identifying a *location* for the metasomatism (the mantle wedge), and more accurately defining what the metasomatising agent *is* (an eclogite melt). Each model shows some major misfits to characteristics of end-member Samoan lavas, although the metasomatic model requires less 'special pleading'.

2. Samples and Methods

The samples analyzed in this study have already been well characterized in terms of major and trace element abundances and isotopic (Sr-Nd-Pb-He-Os) compositions (Workman et al., 2004). Samples selected here are a subset of a much larger collection discussed by Workman et al. (2004) and are used as the result of their containing olivine phenocrysts plentiful enough for separation. Samples with names beginning with a letter (T, U or S) are subaerial samples from the Samoan Islands, and those beginning with numbers are dredged from Vailulu'u and Malumalu Seamounts as well as submarine portions of Ta'u Island.

Whole rock samples were crushed and sieved to obtain a 300-660 μ m size fraction. The sieved fraction was then rinsed in acetone and deionized water. Olivine and clinopyroxene phenocrysts were hand-picked under binocular microscope either dry or from a water bath. The only mineral grains picked for δ^{18} O analyses were those with the fewest visible melt inclusions, mineral inclusions or surface discolorations.

Oxygen isotope compositions ($\delta^{18}O = 1000*(^{18}O/^{16}O_{sample}/^{18}O/^{16}O_{sMOW}) - 1$), where SMOW = standard mean ocean water at an $^{18}O/^{16}O$ ratio of 0.0020052) were determined by laser fluorination (LF) at the Caltech laboratory based on methods described by Valley et al. (1995) and Eiler et al. (2000a). Over 3 days of LF analyses, 16 runs of the GMT-2 garnet show a 1σ reproducibility of 0.07‰ about a mean of 5.83‰ – within error of the accepted, long-term average of 5.80‰ (see Valley, 1995; Eiler et al., 2000a).

Sample runs contain 1-2 mg of multiple mineral grains, and each sample is ideally run in triplicate, although some samples presented here only have 1 or 2 analyses. During LF analyses, some (not all) sample runs showed a high value of the contaminant mass 47 (i.e. \sim 100 times normal). This unknown contaminant was also enriched in mass 48, resulting in highly elevated δ^{18} O values (generally 10-20‰, but up to 37‰) that were all discarded.

This was a problem that was never encountered on a standard run. Consequently, sample splits were heat-treated at 1000° C for 1 and 10 minutes in air, thereby greatly improving the frequency of good runs. Although the heat treatment causes visible oxidation of olivine grains, there is no systematic difference in δ^{18} O between sample splits that were and were not pre-treated; in other words, heat-treated sample splits are both higher and lower than, but within 2σ error of, their non-treated splits with successful LF runs (see Table 1).

The range in δ^{18} O shown by replicate analyses of olivine from a given sample is slightly greater than the quoted reproducibility of the standard. For example, samples 71-2 (n=4) and 78-3 (n=3) both show variations on the order of 0.2‰. Although this is only barely beyond the limit of 2σ reproducibility of the standard, it may be suggestive of minor sample heterogeneity, as shown strongly in a SIMS study of single olivine phenocrysts from Iceland (Gurenko and Chaussidon, 2000) as well as multiple olivine analyses from a single lava flow (also from Iceland; Eiler et al., 2000b).

3. Results

Olivine phenocrysts from 20 Samoan shield lavas have δ^{18} O compositions of 5.11-5.55% (Table 1). The lower end of the Samoan range overlaps with olivines in MORBs and upper mantle peridotites (5.1 \pm 0.2%) that are either measured directly (Mattey, 1994; Eiler et al., 1997) or inferred from melt compositions (Eiler et al., 2000c). No lavas from any other mantle components, including another enriched mantle, EM1 (found at Pitcairn Island; Eiler et al., 1995), have δ^{18} O extending to such high values (see Eiler et al., 1997; Eiler, 2001); exceptions are from lavas that have undergone extensive crystal fractionation (e.g. Muehlenbachs and Byerly, 1982; Geist et al., 1998; Harris et al., 2000) or recent assimilation of altered crustal components (Garcia et al., 1998).

Clinopyroxene (cpx) phenocrysts have been analyzed for δ^{18} O from 5 samples and show a nearly equivalent range in δ^{18} O (4.97-5.53‰) as the olivines. Figure 1 shows that for individual samples, the δ^{18} O compositions of cpx's are similar to or lower than their coexisting olivines. This feature is directly opposite of measured and theoretical equilibrium δ^{18} O partitioning between cpx and olivine, in which cpx is generally 0.4‰ higher than olivine (Chiba et al., 1989; Mattey et al., 1994; Chazot et al., 1997; Macpherson et al., 1998; Harris et al., 2000). From this perspective, if we only had cpx data, the δ^{18} O values found in Samoa would not be an δ^{18} O-enriched anomaly in the global database of oceanic basalts, as the implied δ^{18} O of equilibrium melts would be fully below the upper limit of MORB melts

(basaltic melts are assumed to be 0.5% heavier than olivines and hence 0.1% heavier than cpx; Anderson et al., 1971; Kyser et al., 1981; Eiler, 2001). Although precise knowledge of equilibrium δ^{18} O fractionations as a function of mineral and melt major element compositions, temperature and volatile content are lacking, it is possible that olivine and cpx are seemingly out of equilibrium because they record δ^{18} O at different stages in the evolution of the melts. Because olivine is the first mineral to become saturated in Samoan magmas (along with spinel), we henceforth use olivine as a proxy for δ^{18} O compositions of primary liquids.

Increasing $\delta^{18}O$ values of olivine strongly correlate with increasing incompatible element ratios and heavy isotope compositions (figs. 2-4). Figure 2 shows the relationship between $\delta^{18}O$ and La/Sm, but there are similarly good correlations with other element ratios such as K/Na, K/Ti and Rb/Sr, suggesting that high $\delta^{18}O$ derives from sources that are enriched in the highly incompatible elements. The highest $\delta^{18}O$ occurs in Malumalu lavas (samples 78-1 and 78-3), which also have the highest La/Sm (6.4), $^{208}\text{Pb}/^{204}\text{Pb}$ (39.86) and $^{87}\text{Sr}/^{86}\text{Sr}$ (0.7089). These Malumalu lavas define the most extreme EM2 basalts in the current global database and therefore equilibrium olivine from end-member EM2 is established to have $\delta^{18}O$ equal to 5.5‰. Sources with the lowest $^{87}\text{Sr}/^{86}\text{Sr}$, marked primarily by lavas from Ta'u and Upolu, are almost entirely separate from Malumalu by having lower oxygen and Sr isotopic compositions along with lower La/Sm (figs. 2-4). Vailulu'u olivines show the greatest range in $\delta^{18}O$ of any volcano (~0.4‰), but all other source proxies are nearly identical for the 6 Vailulu'u samples. On the other hand, Malumalu shows the greatest range in $^{87}\text{Sr}/^{86}\text{Sr}$ (0.7067-0.7089) of these volcanoes, but has $\delta^{18}O$ values that vary by only 0.22‰ (fig. 4).

The differences in source characteristics between eastern and western Samoa are highlighted in Figure 3. Western Samoan, comprised of the islands Savai'i and Upolu, overlaps the eastern shield lavas in δ^{18} O but is distinctly lower in 208 Pb/ 204 Pb. Workman et al. (2004) claim that, of all the Samoan volcanoes, Savai'i and Upolu have the least abundant EM2 component and instead are strongly influenced by depleted mantle (for Upolu) and another, yet unidentified, source with low 206 Pb/ 204 Pb and 208 Pb/ 204 Pb but high 207 Pb/ 204 Pb (for Savai'i). The one Savaiian sample presented here is from the oldest volcanic series exposed on the island and yields a δ^{18} O composition (5.38‰) that is mid-range of the whole Samoan suite. Olivine in two young, post-erosional lavas from Savai'i, analyzed by Eiler et al. (1996), have δ^{18} O values (5.48‰ and 5.67‰) extending higher than the end-member EM2 value (5.5‰), but it is unclear if this represents a source signal or is a product

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of shallow assimilation of older, altered Savaiian shield. Since all lavas from Savai'i have similar Sr-Nd-Pb isotopic signatures (Workman et al., 2004), we suggest that the lower δ^{18} O value for Savai'i is primary and the elevated δ^{18} O in post-erosional samples is from secondary processes. No further attempt is made to explain the characteristics of post-erosional lavas, as the primary focus of this work is to determine the origin of high δ^{18} O in the EM2 mantle component.

4. Process Control

The inference of small variations of δ^{18} O in mantle sources based on small variations of δ¹⁸O in mantle-derived lavas must be validated with an assessment of melt generation and fractionation processes. Eiler (2001) shows that joint olivine and plagioclase fractionation, as appropriate for MORBs, results in a nearly negligible net change of δ^{18} O (<0.1%, over a large range of MgO) since light olivine is 'buffered' by heavy plagioclase. However, the Samoan lavas from this study have liquid lines of descent that are saturated with olivine ± clinopyroxene (Workman et al., 2004), minerals that are both lighter than equilibrium melts (see discussion above). A simple calculation, assuming an olivine-melt δ^{18} O fractionation of 0.5‰, indicates that 40% olivine fractionation (an upper limit for magmas presented here) will increase a melt's δ^{18} O composition by 0.2%. Greater increases in a melt's δ^{18} O can only occur if minerals with greater mineral-melt δ^{18} O differences are fractionated. Spinel and chromite are two such minerals (Eiler, 2001) that are present in Samoan lavas, but at a possible fractionated mass of ~2% and mineral-melt δ^{18} O fractionation of 2‰, melt δ^{18} O will increase by less than 0.1%. Fractionation of titanomagnetite has been shown to cause increases in δ^{18} O of silicic Galapagos lavas (Muehlenbachs and Byerly, 1982), but there is no evidence this phase has played a role in Samoan fractionation trends.

An assessment of crystal fractionation effects on melt δ^{18} O values would benefit from more detailed modeling of an AFC-type process (Assimilation-Fractional-Crystallization; DePaolo, 1981) suggested to occur in some OIB lavas. However, an AFC process to create high δ^{18} O may be more important in the formation of EM2 mantle than in the present-day generation of Samoan melts; there is no reason to think that EM2 hotspot chains are unique in the way they generate or process melts, and no other OIB's show elevated δ^{18} O values.

Olivines with higher $\delta^{18}O$ tend to come from whole rocks with lower SiO_2 and CaO than do the lower $\delta^{18}O$ olivines (fig. 5). This negative correlation is slightly deceptive because picritic samples (with low SiO_2 and CaO) are generally from Vailulu'u and

Malumalu (with high $\delta^{18}O$). This observed relationship is not the result of crystal fractionation or accumulation since $\delta^{18}O$ does not correlate with Mg#'s (molar fraction of Mg/[Mg+Fe²⁺]) of whole rocks or of olivines (Jackson et al., unpubl.). Whole rock samples with the lowest Mg#'s are from Upolu, and are the only obvious candidates for showing possible effects of olivine fractionation on $\delta^{18}O$, as other lavas have Mg#'s too high to have fractionated enough olivine to significantly change a melt's $\delta^{18}O$. It is, of course, possible that a melt has experienced crystal fractionation and then accumulated 'foreign' olivine.

The above factors may play in role in causing scatter in plots such as those in Figures 2 through 4, but do not negate the fact that $\delta^{18}O$ values are enriched in olivines from EM2 end-member lavas. Fractionations of $\delta^{18}O$ upon the *generation* of melt from a peridotite lithology will likewise not lead to the observed variation in Samoa's $\delta^{18}O$ values; Eiler (2001) shows that, in theory, the $\delta^{18}O$ composition of a melt is a function of Na₂O content, but the Samoan lavas are within a relatively narrow range of Na₂O (~1-3 wt%) where there is very little leverage on $\delta^{18}O$ (0.1‰). Also, there is no correlation between $\delta^{18}O$ and Na₂O, even if lavas are fractionation-corrected to Mg# 73 by incremental addition or subtraction of equilibrium olivine.

5. Source Control

The correlations of $\delta^{18}O$ (olivine) with trace element and Sr-Pb isotopic enrichment in whole rocks (figs. 2-4) robustly indicate that ^{18}O -enrichment exists in the EM2 source. Eiler et al. (1997) made the same observation with analyses from Savai'i (the post-erosional samples discussed above) and the Society Islands, another EM2 hotspot chain (although not as extreme in EM2 as Samoan lavas). Their interpretation was that EM2 represents a mixture of depleted mantle and 2-6% marine sediment having $\delta^{18}O = 15\%$ and $\delta^{18}O = 15\%$ and $\delta^{18}O = 15\%$ and $\delta^{18}O = 15\%$ and $\delta^{18}O = 15\%$ are over a source's $\delta^{18}O$, as evidenced by high $\delta^{18}O$ found in olivines (5.3-5.5%) and pyroxenes (5.7-5.8%) from metasomatised Savaiian xenoliths. In an earlier study, Hauri et al. (1993) showed that cpx separates from these xenoliths have $\delta^{18}O = 15\%$ in the Samoan lavas; however, the genetic relationship of the xenoliths to the lavas is unclear since Pb isotopic compositions of cpx in the xenoliths do not lay on an extension of trends formed by the lavas (see Workman et al., 2004).

With new light from data presented here and in Workman et al. (2004), it has become clear that the composition of end-member EM2 lavas is much lower in $\delta^{18}O$ and higher in $^{87}Sr/^{86}Sr$ than reported by Eiler et al. (1997). To model the new $\delta^{18}O - ^{87}Sr/^{86}Sr$ trend, sediment having $\delta^{18}O$ of 15-25‰ must have $^{87}Sr/^{86}Sr$ of 0.720 to 0.732 if mixing with primitive mantle and a more narrow range of 0.715 to 0.720 if mixing with depleted mantle; the mass of sediment added would be 2-4%, with less sediment needed as the $\delta^{18}O$ of the sediment increases (see fig. 4). Although the data is roughly fit in these two dimensions, a sediment origin for EM2 is not guaranteed by this calculation alone – there are many more parameters to be checked in order to assess the plausibility of this scenario. Accordingly, in the section below (5.1), we compare a sediment plus mantle mixture to the whole trace element pattern of end-member Samoan lavas.

Marine sediments are not the only material characterized by heavy oxygen. Altered upper oceanic crust (AOC) has been found to have δ 18O values up to ~15-20% (Staudigel et al., 1981, Staudigel et al., 1995; Alt, 2003). When AOC is subducted at convergent margins, elements that are enriched in the ocean crust by low temperature alteration on the seafloor (Rb, Ba, U, K, Pb; Hart and Staudigel, 1989; Staudigel et al., 1996; Bach et al., 2003; Kelley et al., 2003) are also lost from the crust by dehydration and metamorphism, as seen in compositions of eclogites, the high temperature/pressure assemblage of oceanic crust (e.g. Becker et al., 2000). On the contrary, there is evidence from metabasalts and eclogites (Matthews et al., 1984; Neal et al., 1990; Nadeau et al., 1993; Putlitz et al., 2000) that deeply subducted materials do not lose the high $\delta 180$ compositions they acquired from seafloor alteration. Although bulk addition of subducted AOC (i.e. eclogite) would never result in enrichments of the highly-incompatible and light-rare-earth elements, it is possible that a melt derived from such material could have the required trace element enrichments as well as high δ 18O. Below we test a second model for the generation of EM2: metasomatism of the mantle wedge by an eclogite melt derived from a subducting slab. This model expands on the one put forth by Workman et al. (2004) by providing a location (the mantle wedge) and an agent (an eclogite melt) for their proposed mantle metasomatism; however, it is very different in the regard that this is potentially a much more complicated process, involving a greater variety of minerals as well as a greater numbers of 'steps' in the processing of materials.

There are a couple of fairly convincing arguments against a fluid being the enriching component in EM2 mantle: 1) Samoan lavas have slight negative anomalies in the fluid-mobile elements – for example, even though the absolute abundances of U and K are high,

U/Nb and K/La (ratios of 'neighboring' elements in the order of compatibility) are less than in the primitive mantle; and 2) end-member Malumalu lavas have low relative Cl enrichment (see Chapter 4), whereas fluids in subduction zone systems are documented to have high Cl abundances (Kent et al, 2002; Scambelluri et al., 2004).

In summary, the following constraints for EM2, based on isotope and trace elements, must be satisfied in any plausible model for its origin: 1) high time-integrated Rb/Sr and Nd/Sm; 2) elevated δ 18O; 3) lack of significant anomalies across the high-field-strength elements; 4) slight depletions in the fluid-mobile elements.

5.1. Test for a sediment source of enrichment

Present-day assimilation of local marine sediments with high δ^{18} O cannot explain the origin of the Samoan lavas because Pb isotopic compositions of the sediments are too low in ²⁰⁶Pb/²⁰⁴Pb and ²⁰⁸Pb/²⁰⁴Pb to be the enriched component of the end-member Malumalu lavas (see Plank and Langmuir, 1998; Workman et al., 2004). Therefore, we must approach the problem as one of recycling ancient marine sediments back to the mantle at convergent margins, as melting and dehydration processes in the subduction zone have the potential to alter trace element compositions, hence parent/daughter ratios and, consequently over time, radiogenic isotopic compositions. However, difficulties in this problem arise since we have collectively little knowledge about past sediment compositions (that are, in part, a function of the redox state of the oceans), melting and dehydration processes in the subduction zone, and partitioning of trace elements between sediment/fluid and sediment/melt. There is some experimental evidence to suggest that subducted sediments will still look basically like unsubducted sediments (in term of the general sense of anomalies in their trace element patterns), even after dehydration and partial melting, due to sediment/fluid and sediment/melt partition coefficients being close to 1 and often higher (Johnson and Plank, 1999); this does not mean that trace element ratios will not change, but rather that generally enriched or depleted elements will stay generally enriched or depleted, respectively.

There is seemingly no foolproof way of 'selecting' a marine sediment composition to test as an enriching component in EM2, given the many unknowns listed above. Trace elements in sediments that are not thought to fractionate from each other during subduction zone alteration (i.e. Th/La; Plank, 2005) have the potential to fractionate during small-degree mantle melting to make the Samoan lavas. On the other hand, elements that may have similar compatibilities during mantle melting (i.e. Th/U, Ba/Th, K/U, Ce/Pb, Sr/Nd) are likely to

fractionate from each other during dehydration of sediments in subduction zones because of slightly differing fluid mobilities (Johnson and Plank, 1999).

We have chosen to use La/Th and Nb/Th ratios in order to identify a 'test' sediment because these elements are thought to be immobile in sediment-derived fluids and therefore their ratios will not be fractionated by dehydration processes (Johnson and Plank, 1999; Plank, 2005). These ratios show a good correlation with each other for the Samoan lavas (fig. 6) and decent correlations with ⁸⁷Sr/⁸⁶Sr compositions, so must be largely reflective of source variations. We also use Ba/Th as another 'filter' for choosing a sediment; Ba is more fluid-mobile than the other elements used (Johnson and Plank, 1999), but the enormous range of Ba/Th in marine sediments (Plank and Langmuir, 1998) compensates for this possible fractionation in the subduction zone. Using compositions of individual marine sediments from the GLOSS study by Plank and Langmuir (1998), we have found 7 sediments (most of clay-dominated lithology) that lay on an extension of the Samoan trace element trends in La/Th-Nb/Th-Ba/Th space (fig. 6). The average of these 7 selected sediments is reported in Table 2. These 7 out of the 43 sediment compositions reported by Plank and Langmuir (1998) are most distinct from GLOSS by having negative Ba and Sr anomalies.

Figure 7 shows a comparison between trace element patterns of a calculated Malumalu (i.e. EM2) source with mixtures of the selected sediment and depleted or primitive mantle. See Table 2 for details of calculating the Malumalu source composition. Each mixture contains 2% by mass of the sediment and will satisfy the observed $\delta^{18}O$ of Malumalu if the sediment has a δ^{18} O composition of 25‰. To the right of La, the Malumalu source falls between the two sediment-mantle mixtures, except for Pb. To the left of La, the Malumalu source falls below either mixture, except for Nb and Ta, which fall above. Sediment melting is thought to be common beneath arc volcanoes (Elliot et al., 1997; Hoogewerff et al., 1997; Plank and Langmuir, 1998; Johnson and Plank, 1999) and, if it occurred before mixing with the mantle, could potentially account for the excess abundance of Rb, Ba, Th, U and K, but will not be able to explain the low Nb and Ta (Johnson and Plank, 1999), unless a minor phase such as rutile (with high Nb and Ta compatibility; e.g. Rudnick et al., 2000) is present in the sediment during melting. Because of experimental challenges (e.g. Johnson and Plank, 1999), there is very little accurate information about Pb mobility during dehydration and melting. The great enrichment of Pb in the continental crust (42 ppm and 84 times the abundance in primitive upper mantle; Rudnick and Fountain, 1995) must be due to greatly enhanced mobility of Pb at some stage in its generation - whatever that process is could be the same to mobilize Pb from sediments and decrease the high positive anomaly in the sediment-mantle mixtures of Figure 7.

In summary, if sediment recycling to the mantle is the reason for trace element and hence isotopic enrichment in EM2, very specific changes must be made to the sediment before it is mixed and stored with the mantle. While these changes appear unlikely at present, future advancements in the understanding of subduction zone processes may either rule-out or rule-in a 'sediment scenario'.

5.2. Test for an eclogite melt source of enrichment

The trace element composition of eclogite, to be used in the calculation discussed below, is based on eclogite samples with MORB protoliths reported by Becker et al. (2000), and is listed in Table 2. As mentioned above, the addition of bulk eclogite to either primitive or depleted mantle will not result in the required LILE and LREE enrichment in the EM2 reservoir, since these elements are not preferentially enriched over MREE or HREE in eclogites (fig. 8). Even 2 to 3 billion years ago, oceanic crust was most likely not preferentially enriched in these elements, as there is evidence that a depleted upper mantle already existed at that time (Machado et al., 1986; Bowring and Housh, 1995; Vervoort et al., 1996).

Here we test a model in which the mantle wedge (the mantle in-between a subducted slab and the volcanic arc) is metasomatized by a melt from subducted oceanic crust. Recent thermal models for the mantle wedge and subducted slab (van Keken et al., 2002; Kelemen et al., 2003a) suggest that temperatures at the slab-wedge interface may exceed the water-saturated solidus of basaltic compositions beneath the volcanic front, and certainly will at more deeply subducted locations. Melting experiments on garnet pyroxenite show a very narrow temperature range between the solidus and liquidus (Kogiso et al., 2003), meaning melt productivity (as a function of temperature; dF/dT), will be much greater than in peridotite melting.

The upper panel in Figure 8 shows the trace element pattern of our 'test' eclogite. A 30% melt of this eclogite (using partition coefficients in Table 2) is mixed with 1) primitive mantle and 2) depleted mantle. Adding 5% of the melt to each melt-mantle mixture results in trace element patterns that show a fairly decent match to our calculated Malumalu source, especially given the potential errors in the chosen compositions and partition coefficients; to match the δ^{18} O of Malumalu, the eclogite melt must have a δ^{18} O composition of 15‰. The two mixtures bound the Malumalu source for most elements, with the only major misfit

being at U. Since U is a fluid-mobile element, it is possible that eclogites subducted to greater depths will lose additional U by further dehydration reactions.

This scenario has promise in explaining the origin of enrichment in EM2. The caveat if this: the eclogite melt produced from the slab must not be extracted from the mantle wedge, but instead get 'stuck' in it. This may be possible if the melting takes place at subducted depths greater than occur beneath the volcanic arc and the enriched wedge is effectively 'swept away' into greater depths of the mantle.

6. Discussion

Each proposed option for elevated $\delta^{18}O$ in enriched mantle lavas from Samoa requires some form of 'special pleading'. No single scenario can successfully explain the complete array of observation in the lavas, although many fewer discrepancies exist with the model involving metasomatism of the mantle wedge by eclogite melts. Likewise, no single scenario can be confidently ruled out as a possibility due to our limited knowledge base about ancient and present-day subduction zone processing of slabs, as well as compositions of ancient sediments and altered oceanic crust.

If ancient metasomatism by an eclogite melt is a plausible origin for the trace element enrichment in EM2, the reality must be a mixture of sediment and eclogite melts as the enriching component, since the solidus temperatures for these two materials can be very similar (Nichols et al., 1994; Johnson and Plank, 1999; Poli and Schmidt, 2000). This is nearly unavoidable, given the close proximity of sediment overlaying the igneous crust, and the observation that metabasalts and metapelites are often severely intermingled (for example, see Hansteen and Troll, 2003). The exchange of 1% eclogite melt for 0.5% bulk sediment slightly improves the fit of Rb and the Pr-Nd-Sr sedment of the trace element pattern, but makes the negative anomaly in Pb turn positive.

It is possible that some of the second-order variations in the trace element pattern for the Samoan lavas are due to recent alterations to the EM2 source beneath Samoa. For example, recent carbonatite metasomatism has been documented in xenoliths from Savai'i (Hauri et al., 1993). If such a metasomatising material was extracted from the EM2 (Malumalu) source before Malumalu lavas were generated, then the trace element patterns of Malumalu lavas falsely represent their ancient composition. We will need to take great care in understanding the sequence of events in the history of a given source and its derivative melts from the time of their formation until the day of our sampling.

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Figures

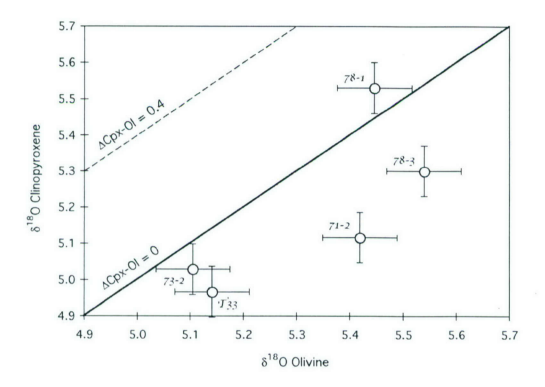


Figure 5-1. Oxygen isotope compositions of coexisting olivine and clinopyroxene from 5 Samoan lavas. Δ Cpx-Ol = δ^{18} O_{Cpx} - δ^{18} O_{Olivine}. Equilibrium Δ Cpx-Ol fractionation is typically reported to be 0.4‰ (Chiba et al., 1989; Chazot et al., 1997; Macpherson et al., 1998; Harris et al., 2000), but the Samoan samples show near zero or negative values. The disequilibrium may be due to olivine and cpx recording magmatic δ^{18} O compositions at different stages in the magmas' evolution. Since olivine is the first phase on the liquidus of Samoan melts, we interpret olivine to be the more reliable recorder of primary magmas. Error bars are plotted as 1σ analytical error (0.07‰).

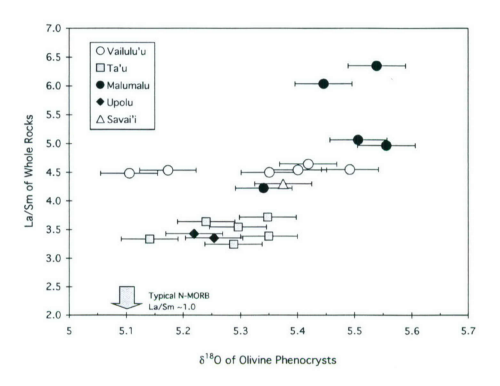


Figure 5-2. Correlation between La/Sm (ppm/ppm) and $\delta^{18}O$ in Samoan lavas. La/Sm ratios can vary as a function of degree of melting, but this is not suspected to control the correlation seen here since La/Sm also correlates well with $^{87}Sr/^{86}Sr$. There are similarly good correlations between $\delta^{18}O$ and other element ratios such as K/Na, K/Ti and Rb/Sr, indicating general enrichment of incompatible elements in sources with elevated $\delta^{18}O$. Error bars are plotted as 1σ standard error (0.05‰) for a sample run in duplicate.

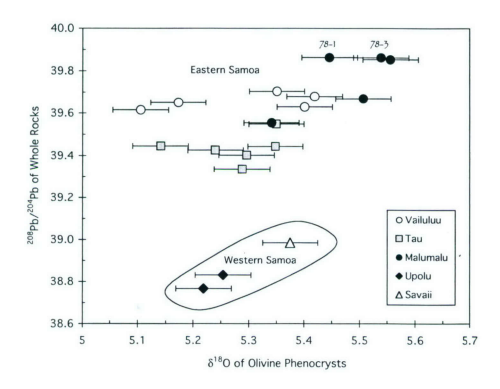


Figure 5-3. Relationship between oxygen and lead isotopes in Samoan lavas. Although the fields for Western Samoa (Savai'i and Upolu) and Eastern Samoa (Vailulu'u, Ta'u and Malumalu) overlap in δ^{18} O values, they are distinct and entirely separate in 208 Pb/ 204 Pb. The source characteristics for Western Samoa lavas are discussed elsewhere (Workman et al., 2004), and the main topic of discussion in this paper is the origin for the extreme mantle enrichment displayed by Malumalu lavas (the two samples with the highest 87 Sr/ 86 Sr in Samoa, 78-1 and 78-3, are identified for reference). Note that there is no Pb isotopic data for the highest δ^{18} O sample from Vailulu'u. Error bars are plotted as 1σ standard error (0.05‰) for a sample run in duplicate.

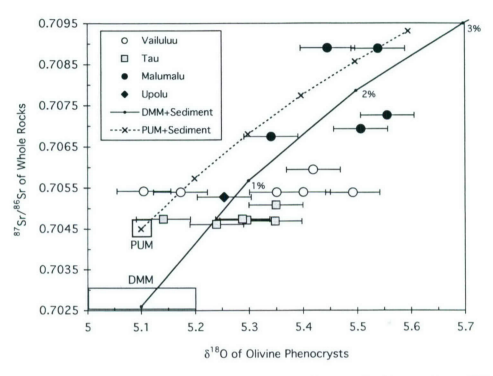


Figure 5-4. Relationship between oxygen and strontium isotopes for Samoan lavas. This positive correlation for enriched mantle lavas has previously been interpreted to represent recycling of sediments from subduction zones back to the mantle (Eiler et al., 1997). Two mixing calculations are shown for reference: Primitive Upper Mantle (PUM) contains 19.9 ppm Sr (McDonough and Sun, 1995), and is mixed with a hypothetical sediment having 170 ppm Sr, 87 Sr, 86 Sr of 0.720 and 18 O of 15‰; Depleted MORB Mantle (DMM) contains 8 ppm Sr (Workman and Hart, 2005) and is mixed with the same sediment except with heavier $δ^{18}$ O (25‰). Tick marks are in increments of 1% sediment addition. Error bars are plotted as 1σ standard error (0.05‰) for a sample run in duplicate.

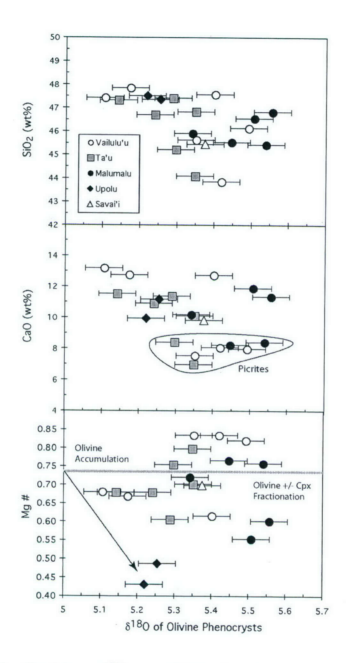


Figure 5-5. Relationships between $\delta^{18}O$ compositions of olivine and whole-rock major element compositions of their host lavas. The line, drawn across the bottom plot at Mg# 73, marks the position of a melt in equilibrium with Fo₉₀ mantle olivine. Lavas above this line are picrites that have accumulated olivine and below this line have fractionated olivine \pm clinopyroxene. The arrow shows the effect on $\delta^{18}O$ from fractional crystallization of 40% olivine. Because there is no correlation between $\delta^{18}O$ and Mg#, $\delta^{18}O$ values are not the result of crystal fractionation.

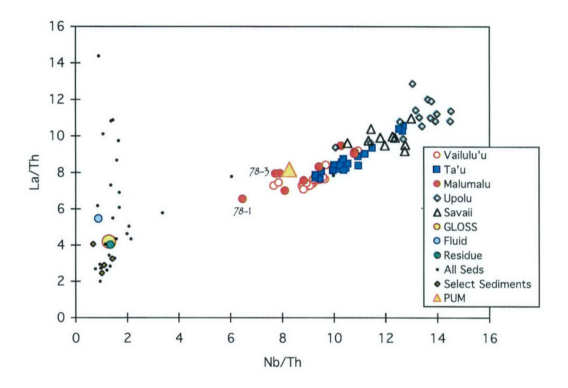


Figure 5-6. Plot of Nb/Th with La/Th ratios of whole rock Samoan basalts (Workman et al., 2004) compared to marine sediments from Plank and Langmuir (1998) used to identify sediment that would be a plausible component of EM2. These ratios were selected because they involve fluid-immobile elements (see Johnson and Plank, 1999; Plank, 2005), so will not be greatly altered by subduction-zone dehydration processes. Since Th is the denominator of each ratio, mixing lines will be linear on this diagram. For reference, DMM has Nb/Th = 18.8 and La/Th = 24.3 (Workman and Hart, 2005). PUM is Primitive Upper Mantle from McDonough and Sun (1995). GLOSS is an average of modern sediments being subducted in trenches today (Plank and Langmuir, 1998). 'All Seds' are individual sediments used to construct GLOSS. 'Select Sediments' are sediments chosen by us to be a plausible mixing component in EM2 based on their position on this plot as well as on a plot of La/Th-Ba/Th. 'Fluid' is a model for slab fluid from Eiler et al. (2000a), and 'Residue' is the residual composition after 7% of the 'Fluid' is removed from GLOSS, indicating insignificant compositional changes to sediment by dehydration (at least for the elements here).

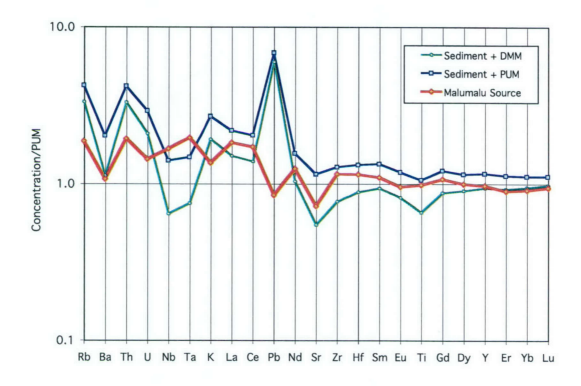


Figure 5-7. Test for mantle enrichment by marine sediments as the origin for EM2. Trace element concentrations are normalized to Primitive Upper Mantle (PUM; McDonough and Sun, 1995). Depleted MORB Mantle is from Workman and Hart (2005). See Figure 6 and Table 2 for sediment selection and derivation of Malumalu source. The Malumalu source is compared with mixtures of sediment with 1) PUM and 2) DMM. The mass fraction of sediment in each mixture is 2%, requiring the sediment to have $\delta^{18}O = 25\%$ to match the $\delta^{18}O$ composition of EM2.

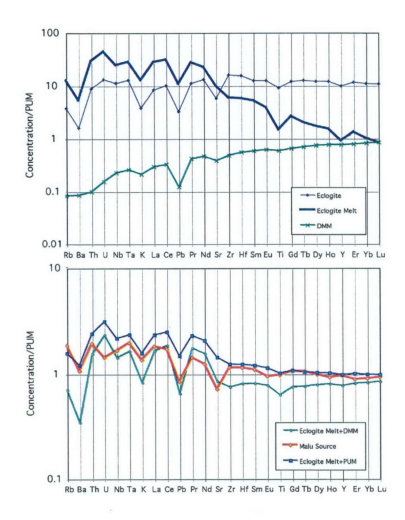


Figure 5-8. Test for metasomatism of depleted/primitive mantle by eclogitic melt as the origin for EM2. Trace element concentrations are normalized to Primitive Upper Mantle (PUM; McDonough and Sun, 1995). Top panel shows 1) the Depleted MORB Mantle (DMM; Workman and Hart, 2005); 2) an average of eclogites with MORB protoliths (Becker et al., 2000; weighted heavily to their sample W1); and 3) a 30% modal, aggregated fractional melt of the eclogite. Bottom panel shows 1) the source for Malumalu lavas assuming that the lavas represent 2.5% aggregated fractional melting; and 2) mixtures of DMM and PUM each with 5% of the eclogitic melt shown in the top panel. The eclogite melt is required to have δ^{18} O of 15‰ to match the δ^{18} O composition of EM2. Trace element compositions, partition coefficients and additional details can be found in Table 2. Because the mantle-melt mixtures show the same general pattern as the calculated Malumalu source (with the only exception being U), and the Malumalu source generally falls in-between the two mantle-melt mixtures (except for the Pr-Nd-Sr segment), we conclude that this scenario is a viable one for the formation of EM2.

Tables

Table 5-1. Oxygen isotope compositions of phenocrysts from Samoan lavas

Average δ^{18} O	5.49	7			5.03	5.03				4.97						5.53	5.30		5.42		
Average	5.49	7														41	4,		4)		
		5.1	5.35	5.40	5.42	5.10	5.35	5.24	5.30	5.14	5.35	5.29	5.34	5.51	5.56	5.45	5.54	5.22		5.25	5.37
					5.531																
					10 min																
			5.348	5.569	5.352				5.336			5.219		5.515	5.54		5.552				
			1 min.	1 min.	1 min.				1 min.			1 min.		1 min.	10 min		1 min.				
			5.316	5.587	5.5	5.07		5.236	5.332	5.166		5.326		5.583	5.641	5.363	5.621				5.376
,			0 min.	0 min.	0 min.	0 min.		1 min.	0 min.	0 min.		0 min.		0 min.	1 min.	0 min.	0 min.				0 min.
	5.492	5.173	5.389	5.381	5.291	5.139	5.348	5.243	5.22	5.115	5.35	5.319	5.341	5.422	5.488	5.529	5.443	5.219		5.254	5.373
	0 min.*	0 min.	0 min.	0 min.	0 min.	0 min.	0 min.	0 min.	0 min.	0 min.	1 min.	0 min.	0 min.	0 min.	0 min.	0 min.	0 min.	0 min.		0 min.	0 min.
cano	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Vailulu'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Ta'u	Malumalu	Malumalu	Malumalu	Malumalu	Malumalu	Upolu	Upolu	Upolu	Savaii
Vol																					
	Volcano	u 0 min.* 5.492	u 0 min.* 5.492 u 0 min. 5.173	u 0 min.* 5.492 u 0 min. 5.173 u 0 min. 5.389 0	u 0 min. * 5.492 u 0 min. 5.173 u 0 min. 5.389 0 u 0 min. 5.381 0	u 0 min.* 5.492 u 0 min. 5.173 u 0 min. 5.389 0 u 0 min. 5.381 0 u 0 min. 5.291 0	u 0 min.* 5.492 u 0 min. 5.173 u 0 min. 5.389 0 u 0 min. 5.291 0 u 0 min. 5.291 0 u 0 min. 5.139 0	0 min. * 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.381 0 min. 5.291 0 min. 5.139 0 min. 5.348	0 min. * 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.139 0 min. 5.57 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.243 1 min. 5.236	0 min. * 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.381 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.348 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236	0 min. * 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.139 0 min. 5.57 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.115 0 min. 5.166	0 min.* 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.139 0 min. 5.57 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.242 0 min. 5.166 1 min. 5.35	0 min.* 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.381 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.115 0 min. 5.166 1 min. 5.35 0 min. 5.319 0 min. 5.326 1 min.	0 min. * 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.243 0 min. 5.07 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.348 0 min. 5.348 0 min. 5.348 0 min. 5.348 0 min. 5.348 0 min. 5.348	0 min.* 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.139 0 min. 5.57 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.115 0 min. 5.166 1 min. 5.35 0 min. 5.319 0 min. 5.326 1 min. 0 min. 5.341 0 min. 5.326 1 min.	0 min. * 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.139 0 min. 5.07 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.35 0 min. 5.35 0 min. 5.35 0 min. 5.341 0 min. 5.341 0 min. 5.383 1 min. 0 min. 5.341 0 min. 5.383 1 min.	0 min. * 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.381 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.115 0 min. 5.166 1 min. 5.319 0 min. 5.326 1 min. 0 min. 5.319 0 min. 5.326 1 min. 0 min. 5.341 0 min. 5.422 0 min. 5.583 1 min. 0 min. 5.422 0 min. 5.583 1 min. 0 min. 5.428 1 min. 5.641 10 min.	0 min. * 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.381 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.243 1 min. 5.27 0 min. 5.243 1 min. 5.236 0 min. 5.348 0 min. 5.343 1 min. 5.336 1 min. 5.35 0 min. 5.349 0 min. 5.336 0 min. 5.349 0 min. 5.356 1 min. 0 min. 5.341 0 min. 5.583 1 min. 0 min. 5.488 1 min. 5.641 10 min. 0 min. 5.443 0 min. 5.363	0 min.* 5.492 0 min. 5.173 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 1 min. 5.35 0 min. 5.319 0 min. 5.326 1 min. 0 min. 5.341 0 min. 5.326 1 min. 0 min. 5.422 0 min. 5.583 1 min. 0 min. 5.488 1 min. 5.641 10 min 0 min. 5.443 0 min. 5.561 1 min. 0 min. 5.443 0 min. 5.621 1 min.	0 min. * 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.381 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.243 0 min. 5.326 1 min. 0 min. 5.341 0 min. 5.326 1 min. 0 min. 5.342 0 min. 5.583 1 min. 0 min. 5.422 0 min. 5.583 1 min. 0 min. 5.443 0 min. 5.641 10 min 0 min. 5.549 0 min. 5.363 0 min. 5.549 0 min. 5.621 1 min.	0 min. * 5.492 0 min. 5.389 0 min. 5.316 1 min. 0 min. 5.291 0 min. 5.587 1 min. 0 min. 5.291 0 min. 5.57 1 min. 0 min. 5.243 1 min. 5.236 0 min. 5.243 1 min. 5.236 0 min. 5.348 0 min. 5.332 1 min. 0 min. 5.349 0 min. 5.336 1 min. 0 min. 5.349 0 min. 5.363 1 min. 0 min. 5.422 0 min. 5.583 1 min. 0 min. 5.488 1 min. 5.641 10 min. 0 min. 5.443 0 min. 5.363 0 min. 5.443 0 min. 5.363 0 min. 5.443 0 min. 5.621 1 min.

^{* 0} min., 1 min. or 10 min. indicate the number of minutes of heat-treatment before analysis of each sample split.

Table 5-2. Compositions and mineral/melt partition coefficients (D's) used in the calculations illustrated by Figures 7 and 8

	Malumalu		Malumalu	Selected			30% Melt of
(mdd)	Lava	Lherzolite D's	Source	Sediment	Eclogite	Eclogite D's	Eclogite
Rb	44.7	0.00002	1.118	97.8	2.27	0.02	7.57
Ba	282.0	0.00012	7.05	344.0	10.54	0.015	35.15
Th	6.17	0.00087	0.154	12.8	669.0	0.015	2.330
n	1.164	0.00101	0.0291	1.978	0.269	0.022	0.895
NP	44.0	0.0035	1.100	13.88	7.39	0.335	16.14
Та	2.92	0.0035	0.0730	0.916	0.477	0.335	1.041
K20 (wt%)	1.574	0.0012	0.0393	2.48	0.110	0.02	0.368
La	44.3	0.009	1.186	39.4	5.44	0.059	18.08
Ce	82.2	0.020	2.875	89.3	16.96	0.139	52.2
Pb	4.35	0.013	0.127	44.1	0.488	0.052	1.62
Pr	9.16	0.026	0.364	1	2.84	0.267	6.97
PN	34.9	0.031	1.558	36.22	16.48	0.495	28.21
Sr	379.1	0.023	14.32	167.8	115.4	0.5	196.1
Zr	205.4	0.046	12.11	154.8	168.3	3.02	62.4
Hf	5.20	0.050	0.325	4.82	4.40	3.02	1.630
Sm	7.15	0.049	0.446	7.25	5.10	2.67	2.122
Eu	1.991	0.061	0.147	1.581	1.938	3.69	0.596
FiO2 (wt%)	2.196	0.078	0.197	0.738	1.844	7.24	0.296
PS	6.28	0.081	0.582	6.23	6.59	5.25	1.443
Tb	0.870	0.109	0.105	1	1.256	7.25	0.201
Dy	4.59	0.135	0.670	5.68	8.22	8.27	1.157
Ho	0.804	0.162	0.139	,	1.812	9.26	0.228
٨	21.89	0.180	4.166	38.46	42.85	12.8	3.93
Er	1.908	0.195	0.392	3.15	5.15	10.2	0.587
Yb	1.399	0.278	0.401	2.94	4.88	12.7	0.450
Lu	0.208	0.297	0.0635	0.449	0.736	15.2	0.057

lithologies, except the heaviest REE are smoothed in garnet Iherzolite and D_{Sr} in eclogite melting is set to 0.5. Spinel Iherzolite inverted source has approximately the correct Rb/Sr, Sm/Nd and Lu/Hf to have developed its observed isotopic composition in 2.5 Ga of aging from PUM (see Workman et al., 2004). The 'Selected Sediment' is the average of 7 sediments from Plank and spinel Iherzolite D's. The Malumalu Source is calculated assuming the lava represents a 2.5% aggregated fractional melt; the composition is based on eclogites with MORB protoliths reported by Becker et al. (2000); the REE elements not analyzed by 1988), and Ta has been calculated assuming Nb/Ta = 15.5. The 30% ecolgite melt assumes aggregated fractional melting of Table 5-2. The Malumalu lava is the average trace element content of samples 78-1 and 78-3, both corrected to Mg# 73 by Langmuir (1998) chosen for their La/Th, Ba/Th and Nb/Th compositions (see fig. 6), and used to test a sediment source of D's are taken from Workman and Hart (2005). The 'Lherzolite D's' represent a mixture of 15% garnet lherzolite and 85% Becker at al. (2000) have been filled-in here by assuming that the REE's will have a slope similar to N-MORB (Hoffman, olivine subtraction. Partition coefficients (D's) are adapted from Kelemen et al. (2003b) for garnet lherzolite and eclogite enrichment for EM2. Note that 3 of the 7 sediments do not have Nb analyses, so are not plotted in Figure 6. The eclogite this eclogite.

Chapter 6:

Gravity-based calculation of crustal flux from the Samoan hotspot and its correlation with Pb-isotopes: a brief overview

Abstract

A Mantle Bouguer Anomaly (MBA) map has been calculated for Samoa from free air gravity and shipboard bathymetry, and then interpreted solely in terms of crustal thickness variations. We find that the western Samoan volcanoes have the greatest amount of excess igneous crust, and that integrated crustal flux decreases nearly monotonically approaching to Vailulu'u Seamount, the easternmost and youngest volcano. This trend shows excellent correspondence to increasing ²⁰⁶Pb/²⁰⁴Pb compositions of the lavas with decreasing age along the hotspot track. We speculate this correlation could be due to either a lithosphere thickening toward the east or decreasing potential temperature of the mantle plume.

1. Introduction

This chapter has been inspired by an age-old question about lavas that are sampled from ocean islands and mid-ocean ridges: are their compositions a product of their source, or a product of the way they are sampled and processed from a multi-component mantle? In other words, is there true spatial heterogeneity in the mantle or are the compositions of surface samples a function of how they are sampled from a 'homogeneously heterogeneous' mantle? The likely story is that both play an important role in magma compositions, but it is unclear how significant each factor could be. The issue of 'source vs. process' is difficult to address with geochemistry alone, as many of the needed physical constraints, such as lithospheric thickness, magma flux, thermal anomalies, and deep mantle structure are largely provided by geophysical observations.

With an extensive geochemical investigation of the Samoan Islands and Seamounts, we have documented systematic changes in lava composition with distance along the hotspot track; this is shown most strongly in the monotonically increasing ²⁰⁶Pb/²⁰⁴Pb compositions with distance from Savai'i toward Vailulu'u Seamount, the present-day hotspot center (see Chapter 2). The goal of this study is to determine whether or not geophysical parameters, specifically crustal flux, will also vary systematically along the volcanic chain.

Using a protocol similar to that of Van Ark and Lin (2004), we have calculated a mantle bouger anomaly map for the Samoan region and interpreted it as variations in total crustal thickness. Integrating excess crustal thickness (total crust minus 6 km of background ocean crust) in cross-sections along the Samoan lineament, we show that crustal flux is systematically decreasing with approach to Vailulu'u. Therefore, increasing ²⁰⁶Pb/²⁰⁴Pb compositions occur as crustal flux wanes. We have not yet developed a detailed model to explain how the geochemical and geophysical properties are linked, but speculate that it may be due to decreasing potential temperature of the Samoan mantle plume or increasing lithospheric thickness over the last 5 million years.

2. Calculations

The calculations described below have been applied to a map region from 168°E to 195°E and 4°S to 16°S within the southern Pacific Ocean (Figure 1). Although we only interpret the results from a small sub-area of this large region, it is as easy to calculate a

large region as it is a small region and we envision the regional scale maps to be used more fully in the future. For example, Hart et al. (2004) have shown that seamounts with a Samoan signature extend back as far as ~175°E, making the oldest known Samoan volcanism ~23 million years old. It would be interesting to compare excess crust and geochemistry in this older section of the Samoan, as we do here for the youngest expression of the Samoan plume.

We employ two kinds of data for this study: free air gravity (FAA) and measured bathymetry. The free air gravity map of Sandwell and Smith (1997) is publicly available at http://topex.ucsd.edu/WWW httml/mar grav.html and was obtained from satellite altimetry measurements taken during the Geosat and ERS1 missions. It has 1-minute by 1-minute resolution and determines the maximum resolution of our calculations since some of the bathymetry data is of higher resolution. The FAA map of our study region is shown in Figure 2. Bathymetry data is taken from two sources: 1) real bathymetry points from the Smith and Sandwell (1997) predicted bathymetry map (also available publicly at: http://topex.ucsd.edu/WWW <a href="http://topex.ucsd.edu/WWW <a href="http://topex.ucsd.edu/WWW httml/mar topo.html) and 2) SeaBeam bathymetry in the region of the Samoan Islands and Seamounts obtained during the 1999 AVON 2 and 3 cruises of the R/V Melville (Hart et al., 2000). The coverage of real bathymetry is shown in Figure 3, with masking of grid nodes that have no data within a 1-minute radius. We have not considered the effect of sediment coverage in our calculations because sediment thicknesses are less than 1000 meters for the entire study region and less than 200 meters for the sub-region involved in our interpretations (fig. 4) (Divins, 2001).

The mantle Bouguer anomaly (MBA) map shown in Figure 5 is calculated, using the Parker (1972) algorithm, by subtracting the attraction of seafloor topography (fig. 3) and a reference model of 6 km thick ocean crust from the free air anomaly (fig. 2). We assume density values of 1.03, 2.7 and 3.3 g/cm³ for seawater, the oceanic crust and the upper mantle, respectively (e.g., Kuo and Forsyth, 1988; Lin et al., 1990). The MBA map and all others to follow have been trimmed by 0.5° on each side in order to edit out the 'edge-effect' of our calculations. Negative values on the MBA map represent regions of excess low-density material (i.e., oceanic crust greater than 6 km); the opposite is true for positive anomalies.

A map of depth to the Moho (base of the oceanic crust), shown in Figure 6, is calculated through a downward continuation of the MBA anomalies to a depth of 11 km (estimated combined depth of 5 km ocean depth plus 6 km reference oceanic crust thickness), as described in greater detail by Van Ark and Lin (2004). This assumes that the entire MBA

signal is due to excesses or deficits in crustal thickness; if there are any mantle thermal effects contributing to the MBA signal, we will have over-estimated depth to the Moho at those locations. The Moho is offset by a constant +3 km in order to force the 'background' Moho depth to be near 11 km, which is the depth of the Moho if the oceanic crust is 6 km thick at locations away from islands or seamounts – for example at 190°E and 12.5°S. Total crustal thickness, shown in Figure 7, is calculated by subtracting the seafloor topography from the calculated Moho depth. Excess crustal thickness is the remaining crust after removing a 6 km thick layer we assume to be 'original' oceanic crust produced at a spreading ridge (fig. 8).

3. Results

Figure 9 show a close-up of excess crustal thickness for the part of the Samoan volcanic chain that is the subject of Chapters 2, 4 and 5; the boundaries of this map are defined by the red box in Figure 1, and coordinates are expressed in kilometers. Although the islands are not plotted here (so as not to obscure the thickness variations), the outlines of islands are clear from the contours of excess crustal thickness (i.e., compare to Figure 1). The calculated maximum excess crust for this area is ~25 km; it occurs beneath Savai'i and is displayed in the cross-section of Savai'i in Figure 9. Moving southeast from Savaii, it is clear from Figure 9 alone that crustal thicknesses are waning. The last volcano of the Samoan chain, Vailulu'u, is shown in cross-section in Figure 9 and has a miximum crustal thickness of ~7 km.

To estimate the crustal flux (i.e., erupted volume flux) from the Samoan plume through time, we have integrated excess crustal thicknesses for 200 km-long cross-sections at every 1 km along the Samoan lineament. Crustal flux along the chain is shown in Figure 10 and is reported as km³ of crust for a 200 km by 1 km map area. Using a Pacific plate speed of 7 cm/yr (Sella et al., 2002), the unit of crustal flux reported in Figure 10 translates to the number of km³ of crust erupted from the Samoan plume in a period of 14,300 years. From Savai'i to Vailulu'u, the calculated crustal flux decreases (almost monotonically except for over the Tutuila-Malumalu ridge) by ~75%. The bottom panel of Figure 10 shows the strong inverse correlation of ²⁰⁶Pb/²⁰⁴Pb to crustal flux. Low ²⁰⁶Pb/²⁰⁴Pb indicates a contribution from trace-element-depleted mantle components to the lavas (i.e., FOZO and DMM) and high ²⁰⁶Pb/²⁰⁴Pb indicates a contribution of trace-element-enriched mantle components to the lavas (i.e., EM2 and HIMU) (see Chapter 2, Figure 16 for correlations

between trace element enrichment and Pb isotopes). Hence, periods of high crustal flux are accompanied by sampling of more depleted mantle components and periods of low crustal flux are accompanied by samples of more enriched mantle components. However, it may be slightly misleading to interpret the crustal thickness variations as if they directly represent temporal changes in magma production from the Samoan plume since recent eruptions have been documented for both Savaii and Vailulu'u (see Chapter 2). Not all the crustal volume from Savaii was produced 5 million years ago (its apparent age from plate motion), but it is unclear what proportion of the crust represents shield versus rejuvenated volcanism.

4. Discussion

The strong relationship between the geochemistry of Samoan lavas and our gravity-based estimate of crustal flux along the chain suggests there is a process-oriented mechanism for producing variations in lava compositions, meaning that lava compositions are a function of what materials are sampled from a multi-component mantle. Below we address two parameters that, if varied systematically, could potentially result in a systematic change in the composition of melts produced from the upwelling plume: lithospheric thickness and potential temperature of the plume.

A very interesting model for melting of a heterogeneous mantle has recently been developed by Ito and Mahoney (2005). They show that the same mantle (composed of multiple components) can produce magmas of varying composition (in trace elements and heavy isotopes) depending on lithospheric thickness at the location of magma genesis. This is due to fundamental differences in the solidus temperatures of the different components and the depth at which melting is 'shut-off' by the lithospheric cap. They assume that, at a given pressure, depleted mantle will melt at a higher temperature than enriched mantle, so melting that is terminated deeply will result in magmas with strong enriched mantle signatures; on the other hand, if the melting column extends to more shallow depths, the enriched melt will be diluted by depleted mantle melts.

Applying this concept to Samoa would suggest that the lithosphere beneath western Samoa is thinner than beneath eastern Samoa, resulting in the production of more melt from the depleted mantle component and, in summation, a greater total melt volume. We will need to explore the implication that western Samoan lavas reflect a greater degree of melting than eastern Samoa lavas; at the present state of investigation, it is not clear if this is the case (as a

side-note, water concentrations in western Samoan lavas may be the key to this question, but such data has not yet been collected).

Changes in the potential temperature of the plume may also explain our observations by controlling the length of the melting column from the bottom instead of the top. At a constant lithospheric thickness, a cooling mantle plume will not be able to produce as much melt from its depleted mantle components, resulting in the geochemical and geophysical evolution from Savai'i to Vailulu'u. However, because of small melt productivity during deep melting (e.g., Asimow et al., 2004), it is possible this scenario will not be able to account for the large differences in crustal volume between western and eastern Samoa.

A changing composition of the Samoan plume (i.e., proportion of depleted to enriched material) would not easily account for coupled variation of calculated igneous flux and ²⁰⁶Pb/²⁰⁴Pb. If the plume changes composition as it upwells beneath the Pacific Plate, but potential temperature and lithospheric thickness remain constant, we would expect the depleted material to produce less total melt volume because of a higher solidus temperature than the enriched material. Our observation is the opposite of this: depleted signatures occur with the greatest crustal volumes.

The last option is the one of coincidence. Consider that the average crustal volume (~1200 km³; Figure 10) over Upolu and Tutuila is the 'constant' crustal volume of a full-grown Samoan shield volcano. Volcanoes to the east have not yet been fully developed, whereas Savai'i has entered a stage of extensive, post-shield, rejuvenated volcanism (as historically documented). From this perspective, geochemical variations would be independent of the igneous flux, unless the average Pb isotopic composition of a volcano changes with growth.

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Figures

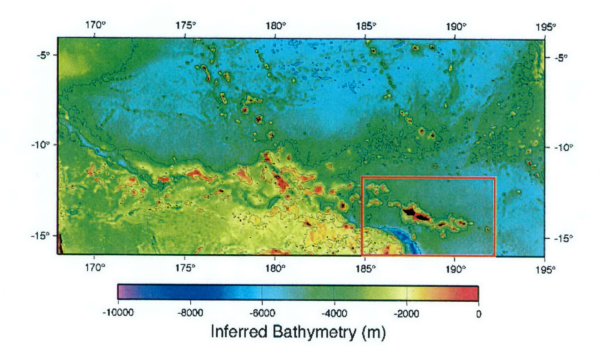


Figure 6-1. Smith and Sandwell (1997) predicted bathymetry for the study region based on a combination of shipboard bathymetric measurements and bathymetry calculated from free air gravity measured by satellite altimetry. The Samoan Islands are shown in black in the southeastern corner of the map. The red box around them marks the boundaries of the map shown in Figure 9.

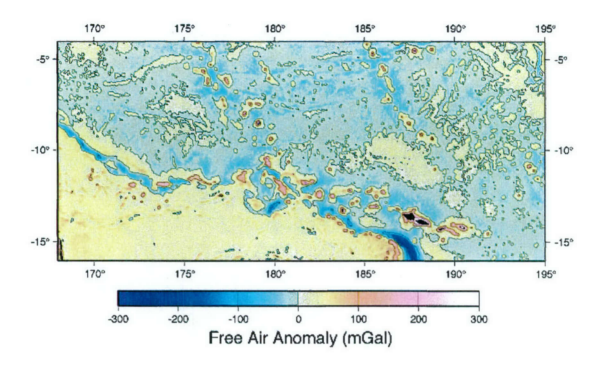


Figure 6-2. Free air anomaly map for the study region from Sandwell and Smith (1997).

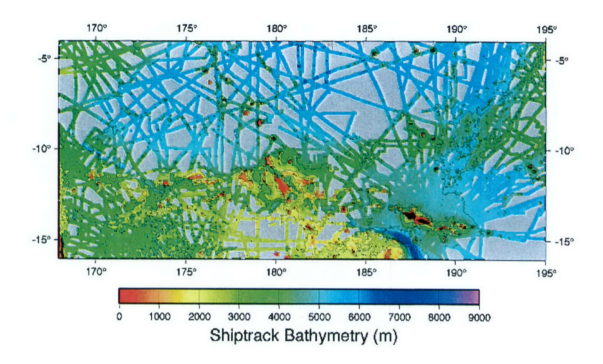


Figure 6-3. Real bathymetry measured by shiptrack. Data sources are Smith and Sandwell (1997) and SeaBeam data from the 1999 AVON 2 and 3 cruises of the R/V/ Melville.

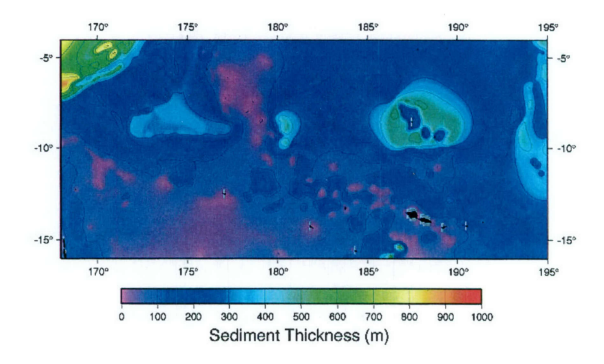


Figure 6-4. Regional sediment thickness map from Divins (2001). Since the sediment cover is generally very thin, we have not considered it in the calculation of MBA.

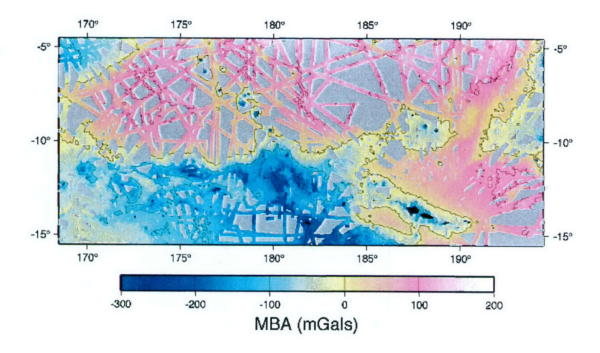


Figure 6-5. Mantle Bouguer anomaly map of the study region calculated according to the Parker (1972) algorithm as implemented by Kuo and Forsyth (1988) and Lin et al. (1990). Negative anomalies indicate excess low-density material, such as over the Samoan Islands and a large portion of the southwest quadrant of the map (some of which may be related to the Samoan plume).

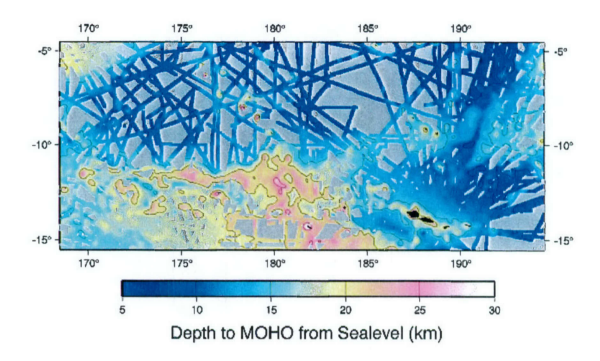


Figure 6-6. Depth to the Moho from sealevel calculated with a downward continuation of the MBA signal, assuming that all the MBA anomaly is due to crustal thickness variations.

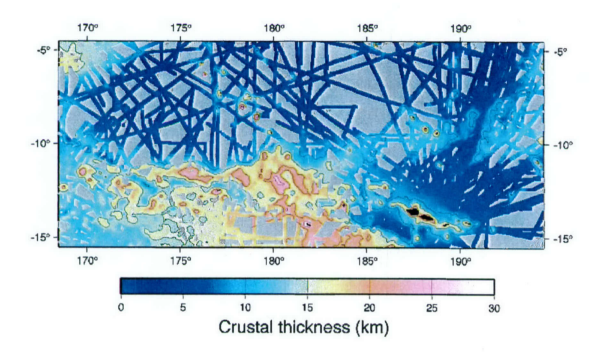


Figure 6-7. Crustal thickness map calculated by subtracting seafloor topography (fig. 3) from the Moho depth (fig. 6).

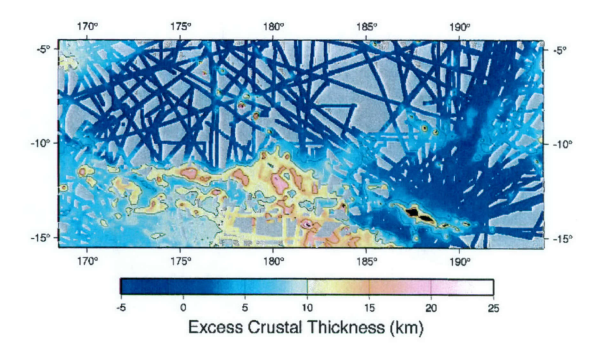
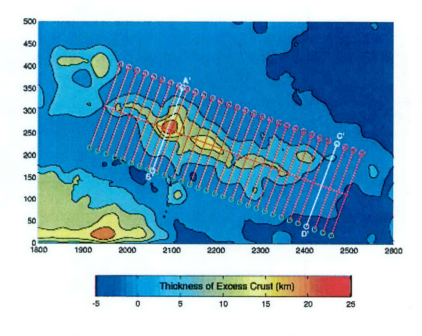


Figure 6-8. Excess crustal thickness calculated by subtracting 6 kilometers from the crustal thickness map in Figure 7. This provides an estimate of crustal flux associated with volcanism after formation of the plate at the East Pacific Rise.



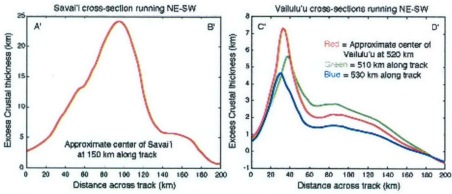


Figure 6-9. Top panel shows excess crustal thickness for the region around Samoa. The boundaries of this map are defined by the red box in Figure 1, but the coordinates here are expressed as kilometers, with the origin being the southwest corner of Figure 1. The 'along-axis' line runs from Savaii at its northwest end to Vailulu'u Seamount at its southeast end. The total distance of the along-axis line is 585 km. 'Across-axis' lines are perpendicular to the along-axis line and are 200 km wide. Plotted here are examples of across-axis lines every 20 km along-axis. Also plotted are two white cross axis lines that mark locations of the cross sections shown in the bottom two panels. Bottom left panel is a cross section of crustal thickness across the middle of Savaii, with a maximum of almost 25 km of crust. At the opposite end is Vailulu'u, shown in the bottom right panel. Maximum excess crustal thickness here is only ~7 km; the two additional cross-sections of Vailulu'u are shown for a semi-3D perspective. For reference, the large mass of excess crust shown at the coordinate (1950, 20) is related to the Tonga Arc, but is not a completely reliable estimate of crustal thickness due to 'edge effects' at this map boundary.

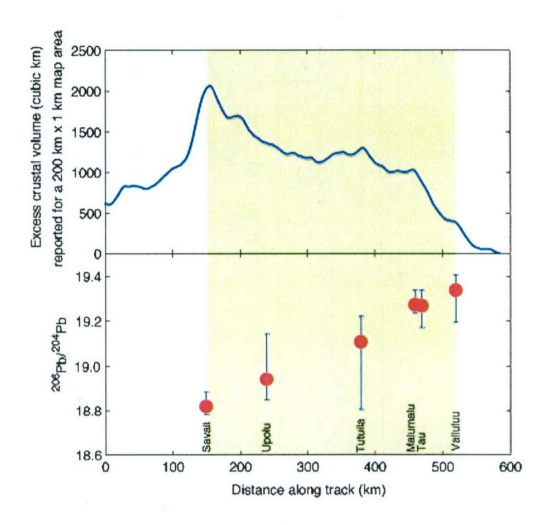


Figure 6-10. A two-panel plot illustrating crustal flux along the Samoan volcanic chain and its surprisingly good inverse correlation with ²⁰⁶Pb/²⁰⁴Pb. Crustal flux is reported as the volume of crust in a 200 km by 1 km area (map view) along the axis shown in Figure 9; i.e., each point is an integration of a cross-section such as those shown in the bottom two panels of Figure 9. Mean ²⁰⁶Pb/²⁰⁴Pb values for each island or seamount are plotted in red, with upper and lower bars indicating the full ranges (data is from Chapter 2). As crustal flux is monotonically decreasing from west to east, mean ²⁰⁶Pb/²⁰⁴Pb is monotonically increasing. We speculate this relationship is due to increasing lithospheric thickness or waning potential temperatures of the Samoan plume over time.

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